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(54) Method for obtaining modified immunoglobulins with reduced immunogenicity of murine antibody variable domains, compositions containing them

Verfahren zur Herstellung modifizierter Immunoglobulinen mit reduzierter Immunogenität der variablen Domänen einer murinen Antikörpers, Zusammensetzungen die diese enthalten

Procédé pour obtenir des immunoglobulines modifiées avec une immunogénicité réduite des domaines variables d'un anticorps murin, compositions les contenant

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(56) References cited:

EP-A-0 519 596 EP-A-0 592 106 EP-A- 0 586 002 WO-A-93/11794

- MOLECULAR IMMUNOLOGY, vol.28, no.4/5, 1, OXFORD, GB pages 489 - 498 E. PADLAN 'A possible procedure for reducing the immunogenicity of antibody variable domains while preserving their ligand-binding properties.'
- PROTEIN ENGINEERING, vol.7, no.6, 4, OXFORD, GB pages 805 - 814 G. STUDNICKA ET AL. 'Human-engineered monoclonal antibodies retain full specific binding activity by preserving non-CDR complementarity-modulating residues.'
- PROTEIN ENGINEERING, vol.4, no.7, 1, OXFORD, GB pages 773 - 783 C. KETTLEBOROUGH ET AL. 'Humanization of a mouse monoclonal antibody by CDR-grafting: the importance of framework residues on loop conformation.
- JOURNAL OF MOLECULAR BIOLOGY, vol.235, no.1, 7 January 1994, LONDON, GB pages 53 - 60 A. CORTI ET AL. 'Idiotope determining regions of a mouse monoclonal antibody and its humanized versions.'

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Description

FIELD OF THE INVENTION

[0001] The present invention is related to the field of immunology, in particular to a method for obtaining modified immunoglobulins with reduced immunogenicity of murine antibody variable domains and compositions containing them.

BACKGROUND OF THE INVENTION

[0002] The immune system builds antibodies that bind to a vast range of antigens with high avidity and specificity, and trigger effector mechanisms. Antibodies have been used in medicine as diagnostic and therapeutic agents, and their potential has been successively enhanced with the advent of new technologies.

[0003] Hybridoma technology allowed isolation of cell lines secreting antibodies of a single specificity (Köhler G., Milstein C. (1975) Nature (London) 256, 495-497), and gene technology has allowed the construction of a range of engineered antibodies from hybridomas.

[0004] Engineering of antibodies is facilitated by their domain structure and may further improve the utility of many antibodies by the acquisition or loss of some of their properties. The antigen-binding properties of the antibody provide the recognition function and this can be attached to one or more of a number of effector agents. The combination of these two features must then be tested against the criteria of efficacy, specificity and immunogenicity.

[0005] Monoclonal antibody producing hybridomas have been most readily obtained from immunized rodents. At present the use of several murine monoclonal antibodies has been widespreaded for the imaging and treatment of malignancy, prophylactic administration to guard against toxic shock, modification of graft rejection episodes, and to temper acute inflammatory reactions.

[0006] In most of the cases where rodent antibodies have been used for therapy, the recipients have elicited an immune response directed towards the antibody. These reactions have limited the duration and effectiveness of the therapy.

[0007] Development of similar reagents from human sources has been frustated, although several options exist, using for example SCID-hu mice, in vitro immunization, recombinatorial libraries, or some useful combination of these. Because there are many well-characterized rodent monoclonal antibodies already available which might be used in the clinic if the immune response could be abolished, the production of engineered antibodies has received much attention.

[0008] Engineered antibodies have been designed to replace as much as possible of the xenogeneic sequences with the equivalent human sequence. Among the genetically engineered antibodies are chimaeric antibodies in which segments from immunoglobulins from diverse species are joined together.

[0009] Initially, chimaeric antibodies were constructed containing the rodent variable regions fused to human constant domains. Particularly mouse/human chimaeric antibodies are potentially useful for immunotherapy for they should exhibit the same specificity but reduced immunogenicity compared to their murine counterparts. The following references describe chimaeric antibody technology: Lobuglio et al, Proc. Natl. Acad. Sci. USA 86: 4220-4224 (1989); United States Patent 4,816,567; PCT International Publication No. WO 87/02671 published May 7, 1987; European Patent Publication No. 255,694 published February 10, 1988; European Patent Publication No.274,394 published July 13, 1988; European Patent Publication No. 323,806 published July 12, 1989; PCT International Publication No. WO 89/00999 published February 9, 1989; European Patent Publication No. 327,000 published August 9, 1989; European Patent Publication No. 328,404 published August 16, 1989; and European patent Publication No. 332,424 published September 12, 1989.

[0010] It is worth noting that even the replacement of the constant regions with human equivalents may not effectively reduce their immunogenicity. Still approximately half of the recipients mounted an immune response to the rodent variable regions. Subsequently, rodent antibodies have been extensively manipulated to resemble more fully human antibodies.

[0011] Further reduction in the immunogenicity of chimaeric antibodies has been achieved by grafting only the complementarity determining regions (CDRs) from the rodent monoclonal antibody onto human framework regions (FRs) prior to its subsequent fusion with an appropriate constant domain (Jones et al, Nature 321: 522-525 (1986)). This procedure to accomplish CDR-grafting often results in imperfectly humanized antibodies, it means, the resultant antibody has either lost affinity or in an attempt to retain its original affinity a number of the murine framework residues have replaced the corresponding ones of the chosen human framework (Winter, European Patent Application, Publication No. 239,400; Riechmann et al, Nature 332: 323-327 (1988)).

[0012] A number of strategies has been developed with the objective of identifying the minimum number of residues for transfer to achieve a useful binding affinity with the least potential consequences on immunogenicity. However, it has emerged that each of these strategies has only been successful to some degree in the reconstitution of parental

affinity.

[0013] The ligand binding characteristics of an antibody combining site are determined primarily by the structure and relative disposition of the CDRs, although some neighbouring framework residues also have been found to be involved in antigen binding (Davies et al, Ann. Rev. Biochem. 59: 439-473 (1990)). Thus, the fine specificity of an antibody can be preserved if its CDR structures and some of the neighbouring residues, their interaction with each other, and their interaction with the rest of the variable domains can be strictly maintained.

[0014] A further procedure for the humanization of an antibody has been suggested by Padlan (Padlan, European Patent Application, Publication No. 0 519 596 A1; Padlan, Molecular Immunology 28: 489-498 (1991)). It is based on the fact that the antigenicity of a protein is dependent on the nature of its surface, and a number of the solvent-accessible residues in the rodent variable region are substituted by residues from a human antibody. The locations of these residues are identified from an inspection of the high resolution X-ray structures of the human antibody KOL and the murine antibody J539. The choice of the human surface residues is arrived at by identifying the most homologous antibody sub-group.

[0015] The nature of the protein surface is important for its recognition and internalization by antigen-processing cells, specifically by antigen-specific B-cells. In addition, the recognition of specific linear sequences by T-cells is also an important element in the immunogenicity of proteins.

[0016] Several groups have developed automated-computerized methods for the identification of sequence features and structural determinants that play a role in the MHC restriction of helper T-cell antigenic peptides (Bersofsky et al, J. Immunol. 138: 2213-2229 (1987), Elliott et al, J. Immunol. 138: 2949-2952 (1987), Reyes et al, J. Biol. Chem. 264: 12854-12858 (1989)). Using these algorithms, it has been possible to identify predicted T cell-presented peptides.

[0017] Analysis of antibodies of known atomic structure has elucidated relationships between the sequence and three-dimensional structure of antibody combining sites (Chothia et al, J. Biol. Chem. 196: 901-917 (1987)). These relationships imply that, except for the third region in the VH domains, binding site loops have one of a small number of main-chain conformations: "Canonical structures". The canonical structure formed in a particular loop is determined by its size and the presence of certain residues at key sites in both the loop and in framework regions.

[0018] An additional subset of framework residues has been defined as "Vernier" zone, which may adjust CDR structure and fine-tune the fit to antigen (Foot et al, J. Mol. Biol. 224: 487-499 (1992)). Substitutions of these residues have been shown to be important to restoring the affinity in CDR grafted antibodies, so the Vernier zone has an obvious consequence for the design of humanized antibodies.

SUMMARY OF THE INVENTION

[0019] It is, accordingly, an objective of the present invention to provide a means of converting a monoclonal antibody of one mammalian species to a monoclonal antibody of another species. Another object is to predict potential T-epitopes within the sequence of variable regions. Another object is to identify the amino acid residues responsible for species specificity or immunogenicity within the sequence of the monoclonal antibody responsible of the T-immunogenicity. Another object is to judiciously replace the amino acid residues within the T-epitope sequences of one species with those of a second species so that the antibodies of the first species will not be immunogenic in the second species. A further object is to make replacements only in the framework regions of the heavy and light chains and not in the complementarity determining regions; also the amino acids belonging to the Vernier zone and those involved in the canonical structures cannot be replaced. Another object of the invention is to provide novel DNA sequences incorporating the replacement amino acid residues. Another object is to provide a vector containing the DNA sequences for the altered antibody. Another object is to provide a eukaryotic or procaryotic host transformed with a vector containing the DNA sequence for the modified antibody.

[0020] A unique method is disclosed for identifying and replacing amino acid residues within T-cell antigenic sequences which converts immunoglobulin antigenicity of a first mammalian species to that of a second mammalian species. The method will simultaneously change immunogenicity and strictly preserve ligand binding properties. A judicious replacement of those amino acid residues within T-cell antigenic sequences of the variable regions, which are not involved in the three-dimensional structure, has no effect on the ligand binding properties but greatly alters immunogenicity.

BRIEF DESCRIPTION OF THE DRAWINGS

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FIGURE 1: Deduced amino acid sequence of (a) VK and (b) VH of murine R3 antibody. CDRs are underlined. FIGURES 2 and 3: Analysis for the modification of the variable regions of heavy and light chains of antibody IOR-R3.

A: sequence of the variable region of the murine IOR-R3 monoclonal antibody.

B: sequence of the variable region of the most homologous human immunoglobulin.

C: sequence of the modified variable region of IOR-R3.

shading: predicted T-cell antigenic sequences, underlined amino acid residues: amino acids involved in tertiary structure.

bold font: complementarity determining regions.

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amino acid residues in boxes: proposed replacements.

The description is the same for both, heavy and light chains.

FIGURE 4: Molecular model of the variable region of mAb R3 displayed as a ribbon. VH is on the right and is darker than VL. The model shows the side chain of murine residues that were mutated in order to humanize the predicted amphipatic segments.

FIGURE 5: Detection of binding of the chimaeric and mutant R3 to EGF-R by RRA.

Antigen binding activity was assayed in different concentrations of purified murine R3 (-.-), chimaeric R3 (+) and mutant VHR3/muR3VK (*) and plotted as CPM of bound ¹²⁵I-EGF against log of the concentration of each antibody. (concentration of IgG was quantitated by ELISA.)

FIGURE 6: Immunization of monkeys with murine R3, chimaeric R3 and mutant R3.

ordinates: Absorbance at 405 nm.

abscises: number of days of blood collected.

The ELISA was performed as described in example 9. The arrows indicate the time of intravenous injection of 2 mg of each mAb. The serum dilution used was 1 / 10 000.

FIGURES 7 and 8: Analysis for the modification of the variable regions of heavy and light chains of antibody IOR-T1.

A: sequence of the variable region of the murine IOR-T1 monoclonal antibody.

B: sequence of the variable region of the most homologous human immunoglobulin.

C: sequence of the modified variable region of IOR-T1 antibody.

The symbols are the same as in FIGURE 2. The description is the same for both, heavy and light chains. FIGURES 9 and 10: Analysis for the modification of the variable regions of heavy and light chains of antibody IOR-CEA1.

A: sequence of the variable region of the murine IOR-CEA1 monoclonal antibody.

B: sequence of the variable region of the most homologous human immunoglobulin.

C: sequence of the modified variable region of IOR-CEA1 antibody.

The symbols are the same as in FIGURE 2. The description is the same for both, heavy and light chains.

DETAILED DESCRIPTION OF THE INVENTION

[0022] The present invention relates to a procedure which simultaneously reduces immunogenicity of the rodent monoclonal antibody while preserving its ligand binding properties in its entirety. Since the antigenicity of an immunoglobulin is dependent on the presence of T-cell antigenic peptides within its sequence, the immunogenicity of a xenogenic or allogenic antibody could be reduced by replacing the residues included in the T-cell antigenic sequences which differ from those usually found in antibodies of another mammalian species.

[0023] The replacement of residues does not include those involved in the canonical structures or in the Vernier zone. This judicious replacement of residues has no effect on the structural determinants or on the interdomain contacts, thus, ligand binding properties should be unaffected as a consequence of alterations which are limited to the variable region framework residues.

(1) Analysis of homology of variable regions

[0024] The present procedure makes use of the available sequence data for human antibody variable domains compiled by Kabat et al, "Sequences of proteins of Immunological Interest", Fifth edition, Bethesda, Maryland; National Inst. of Health, 1994.

[0025] In the first step the variable domains of any heavy or light chain of a first animal species, e.g. the mouse, are

compared with the corresponding variable domains of a second animal species, e.g. human. It is intended that this invention, will allow the antigenic alteration of any animal species antibody.

[0026] The comparison is made by an automated-computerized method (PC-DOS HIBIO PROSIS 06-00, Hitachi). The most homologous human variable regions are then compared, residue for residue, to the corresponding murine regions. This will also define the human subgroup to which each mouse sequence most closely resembles.

(2) Prediction of T-epitopes

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[0027] In the second step, the two homologous variable region sequences, mouse and human, are analysed for the prediction of T-antigenic sequences.

[0028] The algorithm AMPHI (Bersofsky et al, The Journal of Immunology 138: 2213-2229 (1987)) predicts a Helical sequences. The algorithm SOHHA predicts the strip of helix hydrophobicity (Elliott et al, J. Immunol. 138: 2949-2952 (1987)). These algorithms predict T-cell presented fragments of antigenic proteins.

(3) Analysis for immunogenicity reduction

[0029] Those residues in the mouse framework which differ from its human counterpart are replaced by the residues present in the human counterpart. This switching (replacement) occurs only with those residues which are in the T-antigenic sequences.

[0030] Finally, replacement of those residues responsible for the canonical structures or those involved in the Vernier zone could have a significant effect on the tertiary structure. Hence, they cannot be included in the replacement. Additional information about the influence of the proposed replacements on tertiary structure or the binding site could be obtained from a molecular model of the variable regions.

[0031] The molecular model can be built on a Silicon Graphics Iris 4D workstation running UNIX and using the molecular modeling package "QUANTA" (Polygen Corp.).

(4) Method for constructing and expressing the altered antibody

[0032] The following procedures are used to prepare recombinant DNA sequences which incorporate the CDRs of a first mammalian species, usually animal, e.g. murine mAb, both light and heavy chains, into a second mammalian species, preferably human, appearing frameworks that can be used to transfect mammalian cells for the expression of recombinant antibody less immunogenic and with the antigen specificity of the animal monoclonal antibody.

[0033] The present invention further comprises a method for constructing and expressing the modified antibody comprising:

a.-) mutagenesis and assembly of variable region domains including CDRs and FRs regions. The PCR-mutagenesis method (Kamman et al, Nucleic Acids Res. 17: 5404-5409 (1989)) is preferably used to introduce the changes at different positions.

b.-) preparation of an expression vector including one variable region and the corresponding human constant region which upon transfection into cells results in the secretion of protein sufficient for affinity and specificity determinations

c.-) co-transfection of heavy and light chain expression vectors in appropriate cell lines.

[0034] After about 2 weeks, the cell supernatants are analyzed by ELISA for human IgG production. The samples are then analysed by any method for human IgG capable of binding to specific antigens.

[0035] The present invention provides a method for incorporating CDRs from animal monoclonal antibodies into frameworks which appear to be human immunoglobulin in nature so that the resulting recombinant antibody will be either weakly immunogenic or non-immunogenic when administered to humans. Preferably the recombinant immunoglobulins will be recognized as self proteins when administered for therapeutic purpose. This method will render the recombinant antibodies useful as therapeutic agents because they will be either weakly immunogenic or non-immunogenic when administered to humans.

[0036] The invention is further contemplated to include the recombinant conversion of any animal monoclonal antibody into a recombinant human-appearing monoclonal antibody by providing that with a suitable framework region.

[0037] The invention is intended to include the conversion of any animal immunoglobulin to a human-appearing immunoglobulin. It is further intended that human-appearing immunoglobulin can contain either Kappa or Lambda light chains or be one of any of the following heavy chain isotypes (alpha, delta, epsilon, gamma and mu).

[0038] The following examples intend to ilustrate the invention but not to limit the scope of the invention.

EXAMPLE 1: Murine Variable region of R3 monoclonal antibody DNA sequencing

[0039] Cytoplasmic RNA was extracted from about 10⁶ R3 (anti Epidermal growth Factor receptor) hybridoma cells as described by Faloro et al (Faloro, J. et al, Methods in Enzymology 65: 718-749, 1989).

[0040] The cDNA synthesis reaction consisted of 5 ug RNA, 50 mM Tris-HCl, pH 7.5, 75 mM KCl, 10 mM DTT, 3 mM MgCl₂, 25 pmol of CG2AFOR primer (5' GGAAGCTTAGACCGATGGGGCCTGTTGTTTTG 3') for the heavy chain variable region or CK2FOR primer (5' GGAAGCTTGAAGATGGATACAGTTGGTGCAGC 3') for the light chain variable region, 250 uM each of dATP, dTTP, dCTP, dGTP, 15 U ribonuclease inhibitor (RNA guard, Pharmacia) in a total volume of 50 ul. Samples were heated at 70°C for 10 min and slowly cooled to 37°C over a period of 30 min. Then, 100 units MMLV reverse transcriptase (BRL) were added and the incubation at 37°C continued for 1 hour.

[0041] The VH and VK cDNAs were amplified using the PCR as described by Orlandi et al (Orlandi, R. et al, Proc. Natl. Acad. Sci. USA 86: 3833-3837, 1989). For PCR amplification of VH, DNA/primer mixtures consisted of 5 ul cDNA, 25 pmoles of CG2AFOR primer (5' GGAAGCTTAGACCGATGGGGCCTGTTGTTTTG 3') and VH1BACK primer (5' AGGT(G/C)(A/C)A(A/G)CTGCAG(G/C)AGTC(A/T)GG 3').

[0042] For PCR amplification of VK, DNA/primers mixtures consisted of 5 ul cDNA and 25 pmoles of CK2FOR primer (5' GGAAGCTTGAAGATGGATACAGTTGGTGCAGC 3') and VK10BACK primer (5' TTGAATTCCAGTGATGTTTT-GATGACCCA 3'). To these mixtures were added 2.5 mM each of dATP, dCTP, dTTP, and dGTP, 5 ul constituents of 10X buffer thermolase and 1 unit of Thermolase (IBI) in a final volume of 50 ul. Samples were subjected to 25 thermal cycles at 94°C, 30 sec; 50°C, 30 sec; 72°C, 1 min; and a last incubation for 5 min at 72°C. Amplified VH and VK DNA were purified on Prep. A Gene purification kit (BioRad).

[0043] The purified VH and VK cDNA were cloned into M13 vector. Clones were sequenced by the dideoxy method using T7 DNA Pol (Pharmacia). See figure 1.

EXAMPLE 2: Construction of chimaeric genes

[0044] We reamplified the cDNA by PCR using VH1BACK primer (5' AGGT(G/C)(A/C)A(A/G)CTGCAG(G/C)AGTC (A/T)GG 3') and VH1FOR primer (5' TGAGGAGACGGTGACCGTGGTCCCTTGGCCCCAG 3') for VH. and VK3BACK primer (5' GACATTCAGCTGACCCA 3') and VK3FOR primer (5' GTTAGATCTCCAGTTTGGTGCT 3') for VK. The amplified cDNAs were digested with Pstl and BstEll for the VH gene or Pvull and Bglll for the VK gene. The fragments were cloned into M13-VHPCR1 (digested with Pstl and BstEll) or into M13-VKPCR1 (digested with Pvull and Bcll). Details of vectors are given by Orlandi, R. et al, Proc. Natl. Acad. Sci. USA 86: 3833-3837, 1989. The M13VHPCR-R3 and M13VKPCR-R3 containing V gene inserts were identified directly by sequencing.

[0045] The VH gene together with the Ig heavy chain promoter, appropriate splicing sites and signal peptide sequences were excised from M13 vectors by digestion with HindIII and BamHI and cloned into an expression vector (pSVgpt). A human IgG1 constant region (Takahashi, N. et al, Cell 29: 718-749, 1982) was then added as a BamHI fragment. The resultant construction was R3VH-pSVgpt. The construction of the R3VK-pSVhyg was essentially the same except that the gpt gene was replaced by the hygromicin resistance gene and a human Kappa chain constant region was added (Hieter, P.A. et al, Cell 22: 197-207, 1980).

EXAMPLE 3: Modification of the variable domain sequences of IOR-R3 murine monoclonal antibody to humanize the predicted T-cell antigenic sequences

[0046] The variable region sequences of heavy and light chains of R3 were analyzed for T-cell antigenic sequences. It was made by using the computer algorithm AMPHI, which predicts segments of the sequences 11 amino acids in length with an amphipatic helix structure, that is have one side hydrophobic and one side hydrophilic which bind to MHC II molecules.

[0047] Within the variable domain sequence of the heavy chain were predicted 5 segments which are (using Kabat's numbering):

- 1. FR1 between amino acids 3-13.
- 2. FR1 between amino acids 8-20.
- 3. FR2 and CDR2 between amino acids 39-55.
- 4. FR3 between amino acids 74-84.
- 5. FR4 and CDR3 between amino acids 100c-110.

[0048] Figure 2 shows the sequences corresponding to the heavy chain.

[0049] This murine sequence is compared with the immunoglobulin sequences included in the GeneBank and EMBL database. The most homologous human variable region sequence is determined and also the human subgroup to

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which the murine sequence most closely resembles is defined. In this case the human sequence found was a fetal immunoglobulin called HUMIGHVA, which variable region has 75% of homology with the FR regions of the murine immunoglobulin R3.

[0050] Both variable region sequences, human and murine are then compared, residue for residue, and those residues in FR regions which are not involved in the vernier zone or with the canonical structures are selected. Therefore they could be changed by those residues at the same position within the human sequence.

[0051] Finally, this analysis is enriched with computer modeling of the binding site. On the molecular model it is possible to define those replacements which will perturb the tertiary structure of the binding site.

[0052] For the heavy chain of murine R3 we propose 6 replacements:

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- 1. LEU at position 11 by VAL
- 2. VAL at position 12 by LYS

With only these two replacements it is possible to disrupt the amphipatic helix and therefore the predicted T-epitope in the FR1.

- 3. SER at position 75 by THR
- 4. THR at position 76 by SER
- 5. ALA at position 78 by VAL
- 6. THR at position 83 by ARG

[0053] In this case, with the replacements proposed in the FR3, it is humanized.

[0054] The T-cell antigenic sequence in the FR2 contains two PRO which is a very rare amino acid residue in most of the helical antigenic sites, so we propose that it is not a real T-cell epitope.

[0055] In the position 108 at the FR4 appears THR which is present in the same position in some human immunoglobulins, only residue 109 (LEU) is very rare in human, except for this point difference most of the predicted T-cell epitope is human, on this basis it does not need to be modified.

[0056] In Figure 3 the analysis for the light chain of murine R3 is shown.

[0057] In the sequence only one amphipatic helix was predicted, between residue 52-63 corresponding to CDR2 and FR3, and in this region only one point difference exists between murine and human sequences, at position 63. No replacement is proposed, because this murine light chain should be non-immunogenic in human (see molecular modelling).

EXAMPLE 4: Molecular Modelling of mAb R3 VK and VH

[0058] A model of the variable regions of mouse mAb R3 was built using the molecular modeling program QUANTA/ CHARM 4.0 (Molecular Simulations Inc., 1994), running on a 150 MHz Silicon Graphics Indigo Extreme workstation. The VK and VH frameworks were built separately from Fab 26-10 (Jeffrey, P.D et al, Proc. Natl. Acad. Sci. USA 90, 10310, 1993) and Fab 36-71 (Strong, R.K. et al, Biochemistry 30, 3739, 1993), respectively. Fab 26-10 and mAb R3 have 92% homology in the VK frameworks and 88% homology in the whole VK region. The VH frameworks of Fab 36-71 and mAb R3 have 85% homology.

[0059] Coordinates were taken from the Brookhaven Protein Data Bank (entries 1IGI and 6FAB). The frameworks of Fab 36-71 were fitted to the frameworks of Fab 26-10, matching only those residues that have been found to be often involved in the interface between the light and heavy variable regions (Chotia, C. et al, J. Mol. Biol. 186, 651, 1985). The VH domain of Fab 26-10 and the VK domain of Fab 36-71 were then deleted leaving the needed hybrid. Side-chain replacements were performed following the maximum overlap procedure (Snow, M.E. et al, Proteins 1, 267, 1986) and comparing, where possible, with other crystal structures.

[0060] The hypervariable regions of the R3-Variable Light (VL) domain (L1, L2 and L3) were built retaining the same main-chain conformations as in Fab 26-10, since the corresponding CDRs in both antibodies are highly homologous and belong to the same canonical structural groups (Chotia, C. et al, Nature 342, 877, 1989). In the VH domain of mAb R3, CDR H1 belongs to canonical structural group 1, as in Fab 36-71, so the main-chain torsion angles of the parent molecule were kept. CDR H2 corresponds to canonical structural group 2 and the main-chain conformation for this loop was taken from the Fv fragment 4D5 (entry 1FVC), which was selected among other highly resolved structures because of the good matching of its H2 loop base with the framework of Fab 36-71. For all the above mentioned loops comparisons with other CDRs from the Data Bank were made to orient the side chains.

[0061] To model CDR H3, which in mAb R3 was 14 amino acids long, a high temperature molecular dynamics was used for conformational sampling (Bruccoleri, R.E. et al, Biopolymers 29, 1847, 1990). First, the whole structure without CDR H3 was subjected to an energy minimization keeping residues H-94 and H-103 fixed and using harmonic constraints of 10 Kcal/(mole atom A²) for main chain atoms. Then a loop was constructed with an arbitrary conformation starting from the two previously fixed amino acids. Those residues close to the framework were placed taking into

consideration other crystal structures and the top part of the loop was built with an extended conformation avoiding strong steric interactions with the rest of the molecule. For the next modeling steps only CDR H3 and the neighbouring side chains within a distance of 5A⁰ were permitted to move. An energy minimization was first carried out and then a molecular dynamics at 800 K was run for 150 picoseconds. The time step for the run was set to 0.001 picosecond and coordinates were saved every 100 steps. The 120 lowest energy conformations from the dynamics run were extracted and subjected to an energy minimization in which all atoms in the structure were allowed to move. Several low-energy conformations were obtained and the one with the lowest energy was used in the subsequent analyses. Differences between murine and humanized variants of R3 antibody were individually modeled to investigate their possible influence on CDR conformation.

[0062] Amino acid replacements in positions 11, 12 (FR1) and 83 (FR3) in the heavy chain variable region.are quite enough distant from the CDRs-FRs boundaries and should not have any influence on binding affinity. SER 75 residue is pointing to outside, thus the replacement by THR seems not to be important for binding capacity. By contrary THR 76 is accessible from the top of the molecule and could be involved in the interaction with the antigen. But the substitution of THR 76 by SER is a conservative change, leading to no major variations in binding affinity probably.

[0063] The replacement of ALA 78 by VAL should not require steric rearrangements. However VAL 78 could "push" forward ILE 34 (H!). In general, the proposed point mutations should not affect binding affinity according to the computer-aided molecular modelling study (Figure 4).

[0064] The same analysis was done in the light chain variable region of IOR-R3, molecular modelling indicates it is not necessary to make any changes in this region.

EXAMPLE 5: Construction of mutant heavy chain variable region of R3 by PCR mutagenesis

[0065] The changes in the amino acids of mutant heavy chain variable region were constructed using PCR mutagenesis (Kammann, M. et al, Proc. Natl. Acad. Sci. USA 86, 4220-4224, 1989).

[0066] Briefly: Two amplification by PCR: the reaction mixture was: 0.5 ul the VH supernatant of single strand DNA cloned in M13, 25 pmoles mutagenic oligo 1 or 2, 25 pmoles mutagenic oligo 3 or 4 primers (See below the primers sequences). To these mixtures were added 2.5 mM each of dATP, dCTP, dTTP, and dGTP, 5 ul constituents of 10X Vent Polymerase buffer (NEB) and 1 unit of Vent DNA Polymerase (NEB) in a final volume of 50 ul. Samples were subjected to 12-15 thermal cycles at 94°C, 30 sec; 50°C, 30 sec; 75°C, 1 min; and a last incubation for 5 min at 75°C. The products of both PCRs are joined in a second. PCR using the outside primers only (3 and 4). Amplified VH DNA was purified on Prep. A Gene purification kit (BioRad).

[0067] For the changes in the FR1 of LEU 11 and VAL 12 by VAL and LYS, respectively, the following primers were used:

Primer 1: 5' GAAGCCCCAGGCTTCTTCACTTCAGCCCCAGGCTG 3'.

Primer 3: 5' GTAAAACGACGGCCAGT 3'.

[0068] These primers are combined in one PCR.

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Primer 2: 5' CAGCCTGGGGCTGAAGTGAAGAAGCCTGGGGCTTCA 3'.

Primer 4: 5' ACTGGCCGTCGTTTTAC 3'.

[0069] These primers are combined in one PCR.

[0070] Then, the products of both PCRs are combined in one PCR using primers 3 and 4.

[0071] For the changes in the FR3, SER 75, THR 76, VAL 78 and THR 86 by THR, SER, VAL and ARG, respectively, the following primers were designed:

Primer 1: 5' GCAGAGTCCTCAGATCTCAGGCTGCTGAGTTGCATGTAGACTGTGCTGGT-GGATTCGTCTACCGT 3'.

Primer 3: 5' GTAAAACGACGGCCAGT 3'.

[0072] These primers are combined in one PCR.

Primer 2: 5' ACGGTAGACGAATCCACCAGCACAGTCTACATGCAACTCAGCAGCCTGAG-ATCTGAGGACTCTGC 3'.

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Primer 4: 5' ACTGGCCGTCGTTTTAC 3'.

These primers are combined, in one PCR.

[0073] Then, the products of both PCRs are combined in one PCR using primers 3 and 4.

[0074] After mutagenesis VH genes were cloned in expression vectors (pSVgpt) yielding the plasmids R3 mut VHpSVgpt.

EXAMPLE 6: Transfection of DNA into NSO cells

[0075] Four ug of R3VH-pSVgpt and 8 ug R3VK-pSVhyg (chimaeric) or R3 mutant VH-pSVgpt and murine R3VK-pSVhyg were linearized by digestion with Pvul. The DNAs were mixed together, ethanol precipitated and dissolved in 25 ul water. Approximately 10⁷ NSO cells (Rat myeloma NSO is a non-lg secreting cell line) were grown to semiconfluency, harvested by centrifugation and resuspended in 0.5 ml DMEN together with the digested DNA in an electroporation cuvette. After 5 min on ice, the cells were given a single pulse of 170V at 960 uF (Gene-Pulser, Bio-Rad) and left in ice for a further 30 min. The cells were then put into 20 ml DMEN plus 10% fetal calf serum and allowed to recover for 24 hours. At this time the cells were distributed into a 96-well plate and selective medium applied, transfected clones were visible with the naked eyes 14 days later.

EXAMPLE 7: Quantification of IgG production

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[0076] The presence of human antibody in the medium of wells containing transfected clones was measured by ELISA. Microtiter plate wells were coated with goat anti-human IgG (heavy chain specific) antibodies (Sera-Lab). After washing with PBST (phosphate buffered saline containing 0.02% Tween 20, pH 7.5), 20 ul of culture medium diluted in 100 ul of PBST from the wells containing transfectants was added to each microtiter well for 1 hour at 37°C. The wells were then emptied, washed with PBST and either peroxidase-conjugated goat anti human kappa (light chain specific) region antibodies (Sera-Lab) were added and incubated at 37°C for 1 hour, the wells were then emptied, washed with PBST and substrate buffer containing orthophenylenediamine added. Reactions were stopped after a few minutes by the addition of sulphuric acid and absorbance at 492 nm was measured.

40 EXAMPLE 8: EGF Receptor Radioligand Competition assays

[0077] The determination of the affinity constant of the ¹²⁵I-EGF binding to its receptor by murine R3, chimaeric and mutant by rupture of epitopes T antibodies was performed by a homogeneous Radio Receptor Analysis (RRA) with human placenta microsomal fraction (Macias, A. et al, Interferon y Biotecnologia 2: 115-127, 1985).

[0078] These chimaeric and mutant by rupture of epitopes T antibodies were assayed using this technique for its ability to bind to EGF-R (figure 5). Both antibodies bound to EGF-R with the same affinity as the original murine antibody (10-9 M), confirming that the correct mouse variable regions had been cloned and the new antibody isotype did not affect binding. Even more, the changes in the mutant antibody did not affect binding to the antigen.

60 EXAMPLE 9: Immunization of Cercopithecus aethiops monkeys with the murine, chimaeric and VH mutant antibodies

[0079] Three treatment groups with two Cercopithecus aethiops monkeys in each group were immunized with murine R3 mAb, chimaeric R3 antibody and mutant VH R3 antibody, respectively. All the groups were immunized subcutaneously on days 0, 14, 28 and 42, with 2 mg of antibody adsorbed into 5 mg of aluminum hydroxide.

[0080] Blood was collected prior to the first immunization and one week later of each immunization, from all the groups, and the serum was obtained from each sample, and kept at -20°C. The titer of antibodies against the munne R3 mAb was determined by an ELISA technique.

[0081] Costar plates (Inc, high binding) were coated with murine R3 monoclonal antibody at a concentration of 10

ug/ml in bicarbonate buffer (pH 9.6) and incubated overnight. Thereafter, the plates were washed with PBST, were blocked with the same buffer containing 1% BSA during one hour at room temperature.

[0082] The washing step was repeated and 50 ul/well of the different serum dilutions were added. After incubating for 2 hours at 37°C, the plates were washed again and incubated 1 hour at 37°C with alkaline phosphated conjugated goat anti-human total or anti-human IgG Fc region specific antiserum (Sigma, Inc). After washing with PBST the wells were incubated with 50 ul of substrate buffer (1 mg/ml of p-nitrophenylphosphate diluted in diethanolamine buffer (pH 9.8)). Absorbance at 405 nm in an ELISA reader (Organon Teknika, Inc).

[0083] A high IgG response to murine R3 antibody was obtained when this antibody was used as immunogen. A lower but still measurable IgG response (1 / 10 000) to the murine R3 antibody was obtained when monkeys were immunized with the chimaeric antibody, contrary to the results obtained with the mutant Vh version (Figure 6). With the mutant VH R3 antibody no response was measurable after two immunizations, and a small response (1 / 10 000) was measured after 4 immunizations.

EXAMPLE 10: Modification of the variable domain sequences of IOR-T1 murine monoclonal antibody to humanize the predicted T-cell antigenic sequences

[0084] The variable region sequences of heavy and light chains of IOR-T1 were analyzed for T-cell antigenic sequences.

[0085] In the variable domain of the heavy chain 3 segments were predicted, they are:

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- 1. FR1 between amino acids 2-21.
- 2. FR1, CDR1, FR2 between amino acids 29-43.
- 3. FR4, CDR3 between amino acids 97-111.
- ²⁵ [0086] FIGURE 7 shows a comparison with the most homologous human sequence and the replacement proposed, which are 5 at the FR1, 2 at the FR2 and 2 at the FR4.

[0087] The same procedure with the light chain (Figure 8) rendered the following T-cell antigenic segments:

- 1. FR3 between amino acids 60-65.
- 2. FR3, CDR3 between amino acids 79-90.
- 3. CDR3 between aminoacids 93-95A.

[0088] After the analysis we proposed 5 replacement in FR3 at positions: 60, 63, 83, 85 and 87.

35 EXAMPLE 11: Modification of the variable domain sequences of IOR-CEA1 murine monoclonal antibody to humanize the predicted T-cell antigenic sequences

[0089] The variable region sequences of heavy and light chains of IOR-CEA1 were analyzed for T-cell antigenic

- 40 [0090] In the variable domain of the heavy chain two segments were predicted, they are:
 - 1. FR1 between amino acids 1-16.
 - 2. CDR3 and FR4 between residues 96-110.
- [0091] FIGURE 9 shows a comparison with the most homologous human sequence and the replacements proposed, which are 7 at the FR1 and 2 at the FR4.

[0092] The same analysis with the light chain (Figure 10) rendered the following T-cell antigenic segments:

- 1. FR1 between amino acids 1-14.
- 2. CDR2-FR3 between amino acids 55-70.
- 3. FR3-CDR3-FR4 between residues 74-100.

[0093] After the analysis we proposed 4 replacements in FR1 at positions 9, 10, 11 and 13, 11 replacements in FR3 at positions 58, 60, 63, 70, 75, 76, 78, 81, 83, 85 and 87, and 1 replacement in FR4 at position 100.

EXAMPLE 12: Analysis of amphipatic segments in variable regions of immunoglobulin families

[0094] The program AMPHI was included as a subroutine in a program written for reading and processing the im-

munoglobulin sequences from the Kabat Data Base. In processing the sequences the following rearrangements were made:

- Undefined amino acids of type GLX (possible GLN or GLU) were defined as GLN (both GLN and GLU have similar hydrophilicity indexes: -0.22 and -0.64 respectively).
- Undefined amino acids of type ASX (possible ASN or ASP, with hydrophilicity indexes of -0.60 and -0.77) were defined as ASN.
- Other undefined amino acids (empty spaces or "strange" symbols in the sequences were defined as XXX (unknown). The program AMPHI assigns a hydrophilicity value of 0.0 to these amino acids.

[0095] Sequences with more than 5 unknown amino acids (XXX) were not included in the analysis.

[0096] After this preliminary analysis each sequence was processed by the program AMPHI and the results are presented in the form of tables for each immunoglobulin family.

[0097] In tables I to VI the analysis for the six mouse heavy chain families is shown. "Predominant amphipatic regions" (PAR) could be defined at those present in more than 90% of the variable region sequences belonging to each family. For example, comparing the framework one (FR1), a PAR could be defined between the 11 and the 16 amino acid residues for the families I and II, by contrary families III and IV have not amphipatic regions in general from the first amino acid to the 30th. In families V and VI, smaller PARs could be defined from 12-14 and 12-15 residues respectively. [0098] Humanization of the PARs would reduce immunogenicity in patients. The clustering of amphipatic regions in the immunoglobulin variable region frameworks supports the universality of the proposed method, i.e. to humanize these predicted T-cell epitopes by few point mutations.

Claims

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1. A method of modifying an antibody comprising:

comparing the framework amino acids of a variable domain of a first mammalian species with the framework amino acids of variable domains of a second mammalian species;

determining a subgroup of the second mammalian species to which the first mammalian species most closely corresponds;

selecting an antibody from said subgroup whose framework sequence is most similar to the first mammalian species' framework sequence;

identifying amino acid residues of the first mammalian species which differ from the amino acid residues of the selected second mammalian species framework and which are within T-cell antigenic sequences in the variable region of the selected antibody;

selecting the amino acid residues that are not within complementarity determining regions and are not directly involved with canonical structures or Vernier zone;

replacing the amino acid residues in the first mammalian species framework which differ from the amino acid residues of the second mammalian species with the corresponding amino acid residues from the most similar second mammalian species thus identified; and obtaining the modified antibody.

- 2. The method of daim 1 wherein the first mammalian species is mouse.
- 3. The method of claim 1 wherein the second mammalian species is human.
- 4. The method of claim 1 wherein one or more heavy chain constant domains, the light chain constant domain, or both heavy and light chain constant domains of said first mammalian species antibody are replaced by the corresponding constant domain of the second mammalian species antibody.
- 5. The method of claim 1 wherein a modified chimaeric antibody with reduced immunogenicity is obtained from an antibody having a heavy chain variable region shown in Figure 2 sub A and a light chain variable region shown in Figure 3 sub A, by making the following point mutations in the framework regions of the heavy chain:

Heavy chain:

FR1: LEU by VAL at position 11,

VAL by LYS at position 12,
FR3: SER by THR at position 75,
THR by SER at position 76,
ALA by VAL at position 78,
THR by ARG at position 83.

6. The method of claim 1 wherein a modified chimaeric antibody with reduced immunogenicity is obtained from an antibody having a heavy chain variable region shown in Figure 7 sub A and a light chain variable region shown in Figure 8 sub A, by making the following point mutations in the framework regions of the heavy and light chains:

Heavy chain:

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FR 1: LYS by GLN at position 3,
VAL by LEU at position 5,
GLN by GLU at position 6,
LYS by GLN at position 13,
LYS by ARG at position 19,
FR 2: THR by ALA at position 40,
GLU by GLY at position 42,
FR 4: THR by LEU at position 108

FR 4: THR by LEU at position 108, LEU by VAL at position 109,

Light chain:

FR 3: ASP by ALA at position 60, THR by SER at position 63, LEU by PHE at position 83, GLU by VAL at position 85, PHE by TYR at position 87.

7. The method of claim 1 wherein a modified chimaeric antibody with reduced immunogenicity is obtained from an antibody having a heavy chain variable region shown in Figure 9 sub A and a light chain variable region shown in Figure 10 sub A, by making the following point mutations in the framework regions of the heavy and light chains:

Heavy chain:

FR 1: PRO by VAL at position 2, LYS by GLN at position 3, LEU by VAL at position 5, GLU by GLN at position 6, GLY by ALA at position 9, ASP by GLU at position 10, GLU by GLY at position 15; FR 4: THR by LEU at position 108, LEU by VAL at position 109,

Light chain:

FR 1: LYS by SER at position 9,
PHE by THR at position 10,
SER by LEU at position 11,
THR by ALA at position 13,
FR 3: VAL by ILE at position 58,
ASP by SER at position 60,
THR by SER at position 63,
ASP by GLU at position 70,
ILE by VAL at position 75.

SER by ILE at position 76,

VAL by LEU at position 78,
GLN by ASP at position 81,
LEU by PHE at position 83,
GLU by THR at position 85,
PHE by TYR at position 87,
FR 4: ALA by GLN at position 100.

8. A chimaeric antibody having a heavy chain variable region shown in Figure 2 sub A and a light chain variable region shown in Figure 3 sub A, modified by the following point mutations in the framework regions of the heavy chain to reduce immunogenicity:

Heavy chain:

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FR1: LEU by VAL at position 11, VAL by LYS at position 12, FR3: SER by THR at position 75,

THR by SER at position 76, ALA by VAL at position 78, THR by ARG at position 83.

9. A chimaeric antibody having a heavy chain variable region shown in Figure 7 sub A and a light chain variable region shown in Figure 8 sub A, modified by the following point mutations in the framework regions of the heavy and light chains to reduce immunogenicity:

25 Heavy chain:

FR 1: LYS by GLN at position 3,
VAL by LEU at position 5,
GLN by GLU at position 6,
LYS by GLN at position 13,
LYS by ARG at position 19,
FR 2: THR by ALA at position 40,
GLU by GLY at position 42,
FR 4: THR by LEU at position 108,

Light chain:

FR 3: ASP by ALA at position 60, THR by SER at position 63, LEU by PHE at position 83, GLU by VAL at position 85, PHE by TYR at position 87.

LEU by VAL at position 109,

45 10. A chimaeric antibody having a heavy chain variable region shown in Figure 9 sub A and a light chain variable region shown in Figure 10 sub A, modified by the following point mutations in the framework regions of the heavy and light chains to reduce immunogenicity:

Heavy chain:

FR 1: PRO by VAL at position 2, LYS by GLN at position 3, LEU by VAL at position 5, GLU by GLN at position 6, GLY by ALA at position 9, ASP by GLU at position 10, GLU by GLY at position 15;

FR 4: THR by LEU at position 108,

LEU by VAL at position 109,

Light chain:

FR 1: LYS by SER at position 9, 5 PHE by THR at position 10, SER by LEU at position 11, THR by ALA at position 13, VAL by ILE at position 58, FR 3: ASP by SER at position 60, 10 THR by SER at position 63, ASP by GLU at position 70, ILE by VAL at position 75, SER by ILE at position 76, 15 VAL by LEU at position 78, GLN by ASP at position 81, LEU by PHE at position 83, GLU by THR at position 85, PHE by TYR at position 87, FR 4: ALA by GLN at position 100. 20

- 11. A pharmaceutical composition comprising a modified chimaeric antibody as defined in any one of claims 8-10.
- 12. The use of a modified chimaeric antibody as defined in any one of claims 8-10, for the manufacture of a drug directed against tumors.

Patentansprüche

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30 1. Verfahren zum Modifizieren eines Antikörpers, umfassend:

Vergleichen der Framework-Aminosäuren einer variablen Domäne einer ersten Säugetierart mit den Framework-Aminosäuren von variablen Domänen einer zweiten Säugetierart,

Bestimmen einer Untergruppe der zweiten Säugetierart, der die erste Säugetierart am besten entspricht,

Auswählen eines Antikörpers aus dieser Untergruppe, dessen Framework-Sequenz der Framework-Sequenz der ersten Säugetierart am ähnlichsten ist,

Identifizieren der Aminosäurereste der ersten Säugetierart, die sich von den Aminosäureresten des Frameworks der ausgewählten zweiten Säugetierart unterscheiden und die innerhalb der Antigen-Sequenzen für T-Zellen in der variablen Region des ausgewählten Antikörpers liegen,

Auswählen der Aminosäurereste, die nicht in den die Komplementarität bestimmenden Regionen liegen und nicht direkt an den kanonischen Strukturen oder der Vernier-Zone beteiligt sind,

Austauschen der Aminosäurereste im Framework der ersten Säugetierart, die sich von den Aminosäureresten der zweiten Säugetierart unterscheiden, durch die entsprechenden Aminosäurereste der so identifizierten ähnlichsten zweiten Säugetierart und

Gewinnen des modifizierten Antikörpers.

- 2. Verfahren nach Anspruch 1, wobei die erste Säugetierart die Maus ist.
- 55 3. Verfahren nach Anspruch 1, wobei die zweite Säugetierart der Mensch ist.
 - 4. Verfahren nach Anspruch 1, wobei eine oder mehrere konstante Domänen der schweren Kette, die konstante Domäne der leichten Kette oder sowohl konstante Domänen der schweren Kette als auch der leichten Kette des

Antikörpers der ersten Säugetierart durch die entsprechende konstante Domäne des Antikörpers der zweiten Säugetierart ersetzt werden.

5. Verfahren nach Anspruch 1, wobei ein modifizierter chimärer Antikörper mit verminderter Immunogenität aus einem Antikörper mit einer variablen Region der schweren Kette, die in Fig. 2, Teil A gezeigt ist, und einer variablen Region der leichten Kette, die in Fig. 3, Teil A gezeigt ist, erhalten wird, indem folgende Punktmutationen in den Framework-Regionen der schweren Kette vorgenommen werden:

schwere Kette:

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FR1: LEU durch VAL in der Position 11,

VAL durch LYS in der Position 12,

FR3: SER durch THR in der Position 75,

THR durch SER in der Position 76,

ALA durch VAL in der Position 78,

THR durch ARG in der Position 83.

6. Verfahren nach Anspruch 1, wobei ein modifizierter chimärer Antikörper mit verminderter Immunogenität aus einem Antikörper mit einer variablen Region der schweren Kette, die in Fig. 7, Teil A gezeigt ist, und einer variablen Region der leichten Kette, die in Fig. 8, Teil A gezeigt ist, erhalten wird, indem folgende Punktmutationen in den Framework-Regionen der schweren und der leichten Kette vorgenommen werden:

schwere Kette:

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LYS durch GLN in der Position 3,

VAL durch LEU in der Position 5.

GLN durch GLU in der Position 6,

LYS durch GLN in der Position 13,

LYS durch ARG in der Position 19,

FR2: THR durch ALA in der Position 40,

GLU durch GLY in der Position 42,

FR4: THR durch LEU in der Position 108,

LEU durch VAL in der Position 109,

35 leichte Kette:

FR3:

ASP durch ALA in der Position 60,

THR durch SER in der Position 63,

LEU durch PHE in der Position 83,

GLU durch VAL in der Position 85, PHE durch TYR in der Position 87.

7. Verfahren nach Anspruch 1, wobei ein modifizierter chimärer Antikörper mit verminderter Immunogenität aus einem Antikörper mit einer variablen Region der schweren Kette, die in Fig. 9, Teil A gezeigt ist, und einer variablen Region der leichten Kette, die in Fig. 10, Teil A gezeigt ist, erhalten wird, indem folgende Punktmutationen in den Framework-Regionen der schweren und der leichten Kette vorgenommen werden:

schwere Kette:

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FR1: PRO durch VAL in der Position 2,

LYS durch GLN in der Position 3,

LEU durch VAL in der Position 5,

GLU durch GLN in der Position 6, GLY durch ALA in der Position 9,

ASP durch GLU in der Position 10,

GLU durch GLY in der Position 15,

FR4: THR durch LEU in der Position 108,

LEU durch VAL in der Position 109,

leichte Kette:

LYS durch SER in der Position 9, PHE durch THR in der Position 10, SER durch LEU in der Position 11, THR durch ALA in der Position 13, VAL durch ILE in der Position 58, FR3: ASP durch SER in der Position 60, THR durch SER in der Position 63, ASP durch GLU in der Position 70, ILE durch VAL in der Position 75, SER durch ILE in der Position 76, VAL durch LEU in der Position 78, GLN durch ASP in der Position 81, LEU durch PHE in der Position 83, GLU durch THR in der Position 85, PHE durch TYR in der Position 87, ALA durch GLN in der Position 100.

Chimärer Antikörper mit einer variablen Region der schweren Kette, die in Fig. 2, Teil A gezeigt ist, und einer 20 variablen Region der leichten Kette, die in Fig. 3, Teil A gezeigt ist, der mit folgenden Punktmutationen in den Framework-Regionen der schweren Kette modifiziert ist, um die Immunogenität zu vermindern:

schwere Kette:

FR4:

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LEU durch VAL in der Position 11, FR1: VAL durch LYS in der Position 12, SER durch THR in der Position 75, FR3: THR durch SER in der Position 76,

ALA durch VAL in der Position 78, THR durch ARG in der Position 83.

Chimärer Antikörper mit einer variablen Region der schweren Kette, die in Fig. 7, Teil A gezeigt ist, und einer variablen Region der leichten Kette, die in Fig. 8, Teil A gezeigt ist, der mit folgenden Punktmutationen in den Framework-Regionen der schweren und der leichten Kette modifiziert ist, um die Immunogenität zu vermindern:

schwere Kette:

LYS durch GLN in der Position 3, VAL durch LEU in der Position 5, GLN durch GLU in der Position 6, LYS durch GLN in der Position 13. LYS durch ARG in der Position 19. THR durch ALA in der Position 40, FR2: GLU durch GLY in der Position 42,

THR durch LEU in der Position 108, FR4: LEU durch VAL in der Position 109,

leichte Kette:

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ASP durch ALA in der Position 60, THR durch SER in der Position 63. LEU durch PHE in der Position 83, GLU durch VAL in der Position 85, PHE durch TYR in der Position 87.

10. Chimärer Antikörper mit einer variablen Region der schweren Kette, die in Fig. 9, Teil A gezeigt ist, und einer variablen Region der leichten Kette, die in Fig. 10, Teil A gezeigt ist, der mit folgenden Punktmutationen in den

Framework-Regionen der schweren und der leichten Kette modifiziert ist, um die Immunogenität zu vermindern:

schwere Kette:

FR1: PRO durch VAL in der Position 2, LYS durch GLN in der Position 3, LEU durch VAL in der Position 5, GLU durch GLN in der Position 6, GLY durch ALA in der Position 9, ASP durch GLU in der Position 10, GLU durch GLY in der Position 15,

FR4: THR durch LEU in der Position 108, LEU durch VAL in der Position 109.

15 leichte Kette:

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FR1: LYS durch SER in der Position 9, PHE durch THR in der Position 10, SER durch LEU in der Position 11, THR durch ALA in der Position 13, FR3: VAL durch ILE in der Position 58. ASP durch SER in der Position 60, THR durch SER in der Position 63, ASP durch GLU in der Position 70, ILE durch VAL in der Position 75, SER durch ILE in der Position 76, VAL durch LEU in der Position 78, GLN durch ASP in der Position 81, LEU durch PHE in der Position 83, GLU durch THR in der Position 85,

PHE durch TYR in der Position 87, FR4: ALA durch GLN in der Position 100.

Pharmazeutische Zusammensetzung, umfassend einen modifizierten chimären Antikörper nach einem der Ansprüche 8 bis 10.

12. Verwendung eines modifizierten chimären Antikörpers nach einem der Ansprüche 8 bis 10 für die Herstellung eines auf Tumoren gerichteten Medikamentes.

Revendications

1. Procédé pour modifier un anticorps comprenant :

la comparaison des acides aminés structurels d'un domaine variable d'une première espèce de mammifères avec les acides aminés structurels de domaines variables d'une seconde espèce de mammifères;

la détermination d'un sous-groupe de la seconde espèce de mammifères à laquelle correspond de manière la plus proche la première espèce de mammifères;

la sélection d'un anticorps à partir dudit sous-groupe dont la séquence structurelle est la plus similaire à la séquence structurelle de la première espèce de mammifères;

l'identification des résidus d'acides aminés de la première espèce de mammifères qui diffèrent das résidus d'acides aminés de la structure de la seconde espèce de mammifères choisie et qui sont à l'intérieur de séquences antigéniques de cellules T dans la région variable de l'anticorps choisi;

la sélection des résidus d'acides aminés qui ne sont pas à l'intérieur de régions déterminant la complémentarité et qui ne sont pas directement impliqués dans les structures canoniques ou la zone de Vernier ;

le remplacement des résidus d'acides aminés dans la structure de la première espèce de mammifères qui diffèrent des résidus d'acides aminés de la seconde espèce de mammifères par les résidus d'acides aminés correspondants provenant de la seconde espèce de mammifères la plus similaire ainsi identifiée; et

l'obtention de l'anticorps modifié.

- 2. Procédé selon la revendication 1, dans lequel la première espèce de mammifères est la souris.
- Procédé selon la revendication 1, dans lequel la seconde espèce de mammifères est l'être humain.
 - 4. Procédé selon la revendication 1, dans lequel un ou plusieurs domaines constants de chaîne lourde, le domaine constant de chaîne légère ou à la fois les domaines constants de chaîne lourde et légère dudit anticorps de la première espèce de mammifères sont remplacés par le domaine constant correspondant de l'anticorps de la seconde espèce de mammifères.
 - 5. Procédé selon la revendication 1, dans lequel un anticorps chimère modifié avec une immunogénicité réduite est obtenu à partir d'un anticorps ayant une région variable de chaîne lourde représentée sur la figure 2 en A et une région variable de chaîne légère représentée sur la figure 3 en A, en effectuant les mutations ponctuelles suivantes dans les régions structurelles de la chaîne lourde :

Chaîne lourde:

FR 1: LEU par VAL en position 11,

VAL par LYS en position 12,

FR 3: SER par THR en position 75,

THR par SER en position 76,

ALA par VAL en position 78,

THR par ARG en position 83.

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6. Procédé selon la revendication 1, dans lequel un anticorps chimère modifié avec une immunogénicité réduite est obtenu à partir d'un anticorps ayant une région variable de chaîne lourde représentée sur la figure 7 en A et une région variable de chaîne légère représentée sur la figure 8 en A, en effectuant les mutations ponctuelles suivantes dans les régions structurelles des chaînes lourde et légère :

Chaîne lourde : -

FR 1: LYS par GLN en position 3,

VAL par LEU en position 5,

GLN par GLU en position 6,

LYS par GLN en position 13,

LYS par ARG en position 19,

FR 2: THR par ALA en position 40,

GLU par GLY en position 42,

FR 4: THR par LEU en position 108,

LEU par VAL en position 109,

Chaîne légère :

FR 3: ASP par ALA en position 60,

THR par SER en position 63, LEU par PHE en position 83,

GLU par VAL en position 85,

PHE par TYR en position 87.

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7. Procédé selon la revendication 1, dans lequel un anticorps chimère modifié avec une immunogénicité réduite est obtenu à partir d'un anticorps ayant une région variable de chaîne lourde représentée sur la figure 9 en A et une région variable de chaîne légère représentée sur la figure 10 en A, en effectuant les mutations ponctuelles suivantes dans les régions structurelles des chaînes lourde et légère :

Chaîne lourde :

FR 1: PRO par VAL en position 2,

LYS par GLN en position 3, LEU par VAL en position 5, GLU par GLN en position 6, GLY par ALA en position 9, ASP par GLU en position 10, GLU par GLY en position 15, THR par LEU en position 108, LEU par VAL en position 109,

Chaîne légère :

FR 4:

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LYS par SER en position 9, FR 1: PHE par THR en position 10, SER par LEU en position 11, THR par ALA en position 13, VAL par ILE en position 58, FR 3: ASP par SER en position 60, THR par SER en position 63, ASP par GLU en position 70, ILE par VAL en position 75, SER par ILE en position 76, VAL par LEU en position 78, GLN par ASP en position 81, LEU par PHE en position 83, GLU par THR en position 85,

8. Anticorps chimère ayant une région variable de chaîne lourde représentée sur la figure 2 en A et une région variable de chaîne légère représentée sur la figure 3 en A, modifié par les mutations ponctuelles suivantes dans 30 les régions structurelles de la chaîne lourde pour réduire l'immunogénicité :

Chaîne lourde:

FR 4:

LEU par VAL en position 11, FR 1:

VAL par LYS en position 12,

PHE par TYR en position 87, ALA par GLN en position 100.

SER par THR en position 75, FR 3:

THR par SER en position 76, ALA par VAL en position 78, THR par ARG en position 83.

9. Anticorps chimère ayant une région variable de chaîne lourde représentée sur la figure 7 en A et une région variable de chaîne légère représentée sur la figure 8 en A, modifié par les mutations ponctuelles suivantes dans les régions structurelles des chaînes lourde et légère pour réduire l'immunogénicité :

Chaîne lourde:

LYS par GLN en position 3, FR 1:

VAL par LEU en position 5,

GLN par GLU en position 6,

LYS par GLN en position 13,

LYS par ARG en position 19,

THR par ALA en position 40, FR 2:

GLU par GLY en position 42,

THR par LEU en position 108, FR 4:

LEU par VAL en position 109,

Chaîne légère :

ASP par ALA en position 60, FR 3: THR par SER en position 63, LEU par PHE en position 83, GLU par VAL en position 85, PHE par TYR en position 87.

10. Anticorps chimère ayant une région variable de chaîne lourde représentée sur la figure 9 en A et une région variable de chaîne légère représentée sur la figure 10 en A, modifié par les mutations ponctuelles suivantes dans les régions structurelles des chaînes lourde et légère pour réduire l'immunogénicité :

Chaîne lourde :

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PRO par VAL en position 2, FR 1: LYS par GLN en position 3, LEU par VAL en position 5, GLU par GLN en position 6, GLY par ALA en position 9, ASP par GLU en position 10, GLU par GLY en position 15, THR par LEU en position 108, FR 4:

LEU par VAL en position 109,

Chaîne légère :

LYS par SER en position 9, FR 1: PHE par THR en position 10, SER par LEU en position 11, THR par ALA en position 13, VAL par ILE en position 58, FR 3: ASP par SER en position 60, THR par SER en position 63, ASP par GLU en position 70, ILE par VAL en position 75,

> VAL par LEU en position 78, GLN par ASP en position 81, LEU par PHE en position 83, GLU par THR en position 85, PHE par TYR en position 87, ALA par GLN en position 100.

SER par ILE en position 76,

- 11. Composition pharmaceutique comprenant un anticorps chimère modifié selon l'une quelconque des revendications 8 à 10.
- 12. Utilisation d'un anticorps chimère modifié selon l'une quelconque des revendications 8 à 10, destinée à la fabri-45 cation d'un médicament ciblant les tumeurs.

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FIGURE 1: DEDUCED AMINO ACID SEQUENCES

A VK OF MURINE R3 ANTIBODY

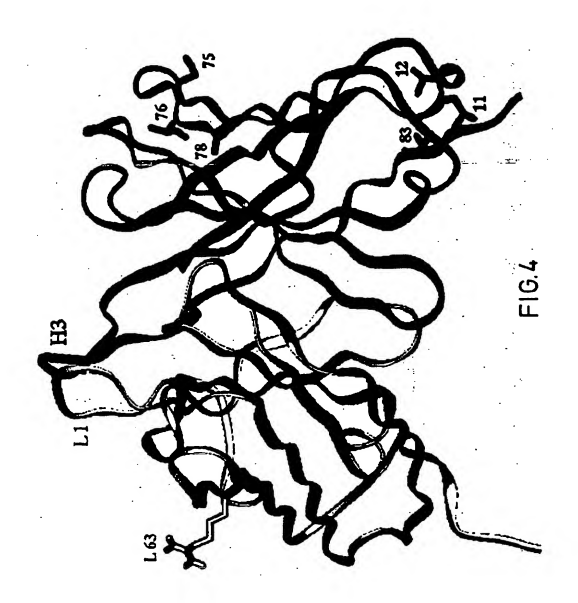
D V L M T Q I P L S L P V S L G D Q A S I S C R S S Q N I N I V H S N G N T Y L D W Y L Q K P G Q S P N L L I Y K V S N R F S G V P D R F R G S G S G T D F T L K I S R V E A E D L G V Y Y C F O X S H V P W T F G G G T K L E I K R A

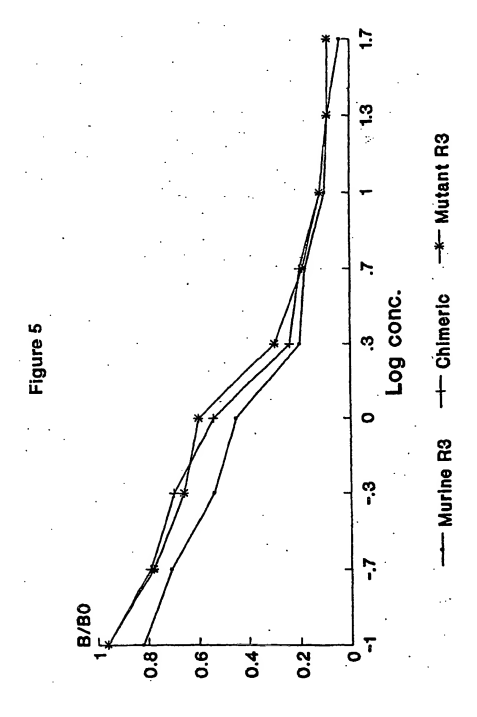
B VH OF MURINE R3 ANTIBODY

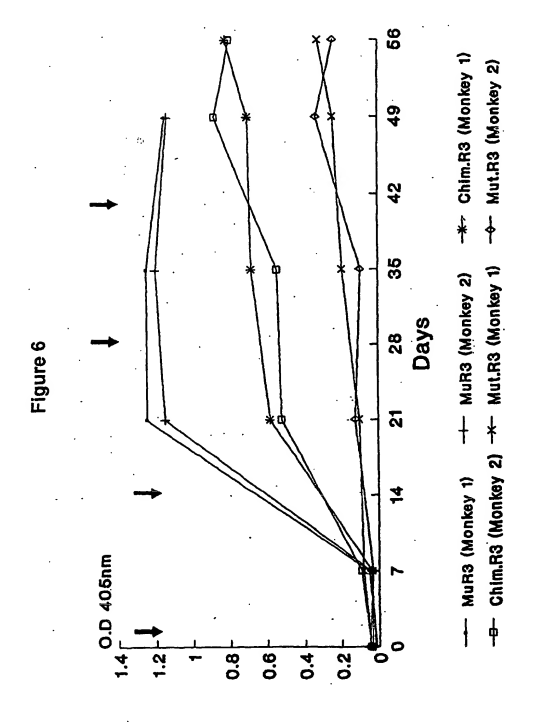
Q V Q L Q Q P G A E L V K P G A S V K L S C K A S G Y
T F T N Y Y I Y W V K Q R P G Q G L E W I G G I N P T
S G G S N F N E K F K T K A T L T V D E S S T T A Y M
Q L S S L T S E D S A V Y Y C T R Q G L W F D S D G R
G F D F W G Q G T T L T V S S

		FIGUE	RE 2: VA	RIABLE	REGIO	N OF T	Æ HEAV	Y CHAI	N OF IC	R-R3.	1	
	1	-	2	A	5	6	7	8	9	10 .	11 ′	12
Α	GLN	VAL	GLN	LEU.		GLN	PRO	GLY	ALA	GLU	LEU	
В	GLN	VAL	GLN	LEU	VAL	GLN	SER'.	GLY	ALA	GLU	VAL	LYS
C·	GLN	VAL	GLN	LEU	GLN	GLN	PRO	GLY	ALA	GLU	VAL	LYS
			•		•							
•	13	14	15	16	17	18	. 19	20	21	22	23	24
A	LYS	PRO	GLY	ALA	SER	VAL	LYS	LEU	SER	CYS	LYS	ALA
В	LYS	PRO	GLY	ALA	SER	VAL	LYS	VAL	SER		LYS	ALA
C	LYS	PRO	GLY	ALA	SER	VAL	LYS .	LEU	SER	CYS	LYS	ALA
						70	33 .	22	33	34	35	36
	- 25	26	27	28	29 .	30	31 ·	32 TVD		<u>ill</u>	TYB	TRP
A	SER	<u>GLY</u>	TYR	THR	PHE	THR	ash	TYR	1111	. IVE	110	TRP
В.	SER	GLY	TYR	THR	PHE	ASN	ASH	TYB	TYR .	<u>ILE</u>	TYR	TRP
C	SER	GLY .	TYR	THR	PHE	THR	MON	110	1 (13	125	* * * * * * * * * * * * * * * * * * * *	110
	37	·38	39	40	41	42	43	44	45	46	47	48
Α	. VAL		GLN		PRO				JEU.		TRP	TIPE)
В	VAL	ARG	GLN	ALA	PRO	GLY	GLN	GLY	LEU	GLU	TRP	MET
Č	: VAL	LYS	GLN	ARG	PRO	GLY	GLN	GLY	LEU	GLU	TRP	ILE
•									•			
	49	50	51	52	52A	53	54	35	56	57	58	59
Α	GLY	CLY	III.	ASH /	PRO	THR	SER	GLY	ELY	SER	KZA	PHE
В	GLY								61 V	OFR	acu.	BUP .
, C	GLY	GLY	ILE	ash	<u>PRO</u>	THR	<u>ser</u>	<u>gly</u>	GLY .	SER	ash	PHE
•				<i>(</i> 2	i.	<i>(</i> =		67	68	69	70	71 .
	60	61 GLU	62	63 Phe	64 LYS	65 Thr	66 LYS	ALA	THR	LEU	THR	VAL
Α	asn	324 34										
		610	LYS	1 116		• • • • • • • • • • • • • • • • • • • •				MET		
В	•						ARG	VAL	THR	MET	THR	ARG
	ASH	CTA	LYS	PHE	LYS	THR				MET		
В	ASH	CTA	LYS .	PHE	tys ·	THR	ARG LYS	VAL ALA	THR THR		THR	ARG
B C	ASH 72	G18 73	1YS	PHE 75	LYS	THB 77	ARG LYS 78	VAL	THR THR	LEU 81	THR THR 82	ARG VAL
B C	ASH 72 ASP	GLU 73 GLU	LYS .	PHE 75	LYS	THB 77	ARG LYS 78	VAL ALA 79	THR THR	LEU 81	THR THR 82	ARG VAL 82A
B C A B	ASH 72	G18 73	LYS 74 ESER	PHE 75 SER	IYS 76 THR	THR 77 THR	ARG LYS 78 ALA	VAL ALA 79	THR THR 80	LEU 81 GLAV	THR THR 82	ARG VAL 82A SER
B C	ASH 72 ASP ASP	GLU 73 GLU THR	LYS 74 SER SER	PHE 75 SER THR	1YS 76 THR SER	THR 77 LHR THR	ARG LYS 78 ALA VAL VAL	VAL ALA 79 TYR	THR THR 80 MET MET	81 GLV GLU GLN	THR THR 82 LEU LEU LEU	ARG VAL 82A SER SER SER
B C A B	ASH 72 ASP ASP	GLU 73 GLU THR GLU 82C	1YS 74 SER SER SER SER 83	PHE 75 SER THR THR 84	1YS 76 THR SER SER	THR 77 IHE THR THR THR 86	ARG LYS 78 ALA: VAL VAL 87	VAL ALA 79 TYR TYR TYR 88	THR THR 80 MET MET 89	81 GLN GLU GLN 90	THR THR 82 LEU LEU 91	82A SER SER SER 92
B C A B	ASH 72 ASP ASP ASP	GLU 73 GLU THR GLU 82C	1YS 74 SER SER SER SER SER	PHE 75 SER THR THR 84 SER	76 THR SER SER GLU	THR 77 IMR THR THR THR 86 ASP	ARG LYS 78 ALA: VAL VAL VAL 87 SER	VAL ALA 79 TYR TYR TYR 88 ALA	THR THR 80 MET MET 89 VAL	81 GLN GLN GLN 90 TYR	THR THR 82 LEU LEU 191 TYR	82A SER SER SER SER SER CYS
B C A B C	ASH 72 ASP ASP ASP ASP	GLU 73 GLU THR GLU 82C LEU	1YS 74 SER SER SER SER TURE	PHE 75 SER THR THR 84 SER SER	76 THR SER SER GLU GLU	THR 77 THR THR THR ASP ASP	ARG LYS 78 ALA VAL VAL 87 SER THR	VAL ALA 79 TYR TYR BB ALA ALA	THR THR 80 MET MET MET VAL VAL	81 GLW GLW GLN 90 TYR TYR	THR THR 82 LEU LEU LEU 71 TYR TYR	82A SER SER SER CYS CYS
B C A B C	ASH 72 ASP ASP ASP ASP	GLU 73 GLU THR GLU 82C	1YS 74 SER SER SER SER SER	PHE 75 SER THR THR 84 SER	76 THR SER SER GLU	THR 77 IMR THR THR THR 86 ASP	ARG LYS 78 ALA: VAL VAL VAL 87 SER	VAL ALA 79 TYR TYR TYR 88 ALA	THR THR 80 MET MET 89 VAL	81 GLN GLN GLN 90 TYR	THR THR 82 LEU LEU 191 TYR	82A SER SER SER SER SER CYS
B C A B C	ASH 72 ASP ASP ASP SER SER SER	GLU 73 GLU THR GLU 82C LEU LEU	14 SER SER SER SER ARG ARG	75 SER THR THR 84 SER SER SER	76 FHR SER SER SER GLU GLU GLU	THR 77 THR THR THR 86 ASP ASP ASP	78 ALS VAL VAL 87 SER THR SER	VAL ALA 79 TYR TYR 88 ALA ALA ALA	THR THR 80 MET MET MET VAL VAL	81 GLN GLU GLN 90 TYR TYR TYR	THR THR 82 LEU LEU 91 TYR TYR	82A SER SER SER CYS CYS CYS
B C A B C	ASH 72 ASP ASP ASP SER SER SER SER	GLU 73 GLU THR GLU 82C LEU LEU 94	14 SER SER SER SER ARG ARG	PHE 75 SERT THR THR 84 SERT SER SER 96	76 FHR SER SER SER GLU GLU 97	THR 77 LHR THR THR ASP ASP ASP 98	78 ALA VAL VAL 87 SER THR SER 99	VAL ALA 79 TYR TYR TYR 88 ALA ALA ALA 100	THR THR 80 NET MET MET VAL VAL VAL	81 GLN GLU GLN 90 TYR TYR TYR 100B	THR THR 82 LEU LEU 191 TYR TYR TYR 100C	82A SER SER SER CYS CYS CYS CYS
B C A B C A	ASH 72 ASP ASP ASP SER SER SER SER 93 THR	GLU 73 GLU THR GLU 82C LEU LEU LEU 94 ARG	14 SER SER SER SER ARG ARG	75 SER THR THR 84 SER SER SER	76 FHR SER SER SER GLU GLU GLU	THR 77 THR THR THR 86 ASP ASP ASP	78 ALS VAL VAL 87 SER THR SER	VAL ALA 79 TYR TYR 88 ALA ALA ALA	THR THR 80 MET MET MET VAL VAL	81 GLN GLU GLN 90 TYR TYR TYR	THR THR 82 LEU LEU 191 TYR TYR TYR 100C	82A SER SER SER CYS CYS CYS
B C A B C A B	ASH 72 ASP ASP ASP SER SER SER SER STHR ALA	GLU 73 GLU THR GLU 82C LEU LEU LEU 94 ARG ARG	LYS 74 SER SER SER 83 FUR ARG ARG 95 CLN	PHE 75 SER*: THR THR 84 SER: SER SER 96 GLY	76 FHR. SER SER SER GLU GLU 97 LEU	THR 77 THR THR THR ASP ASP ASP ASP TRP	ARG LYS 78 ALA VAL VAL SER THR SER 99 PHE	79 TYR TYR 88 ALA ALA ALA 100 ASP	THR. THR 80 MET MET MET VAL VAL 100A SER	81 GLU GLU GLN 70 TYR TYR TYR 100B	THR THR 82 LEU LEU 91 TYR TYR TYR 100C	82A SER SER SER CYS CYS CYS CYS 100D
B C A B C A	ASH 72 ASP ASP ASP SER SER SER SER 93 THR	GLU 73 GLU THR GLU 82C LEU LEU LEU 94 ARG	14 SER SER SER SER ARG ARG	PHE 75 SERT THR THR 84 SERT SER SER 96	76 FHR SER SER SER GLU GLU 97	THR 77 LHR THR THR ASP ASP ASP 98	78 ALA VAL VAL 87 SER THR SER 99	VAL ALA 79 TYR TYR TYR 88 ALA ALA ALA 100	THR THR 80 NET MET MET VAL VAL VAL	81 GLN GLU GLN 90 TYR TYR TYR 100B	THR THR 82 LEU LEU 191 TYR TYR TYR 100C	82A SER SER SER CYS CYS CYS CYS
B C A B C A B	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C LEU LEU LEU 94 ARG ARG ARG	14 SER SER SER SER ARG ARG GIN GIN	PHE 75 SER*: THR THR 84 SER: SER SER 96 GLY GLY	76 FHR. SER SER SER B5 GLU GLU GLU 97 LEU	THR 77 THR THR THR ASP ASP ASP ASP TBP	78 ALA VAL VAL SER THR SER 99 PHE	79 TYR TYR 88 ALA ALA ALA 100 ASP	THR THR 80 MET MET MET VAL VAL VAL SER SEB	81 GLU GLN 90 TYR TYR TYR TYR ASP	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GIV	82A SER SER SER CYS CYS CYS CYS 100D ARG
B C A B C A B C	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C UEU LEU LEU 94 ARG ARG ARG	14 SER SER SER SER ARG ARG ARG GLN GLN	PHE 75 SER*: THR THR 84 SER: SER SER 96 GLY GLY	76 FHR. SER SER SER GLU GLU 97 LEU 103-	THR 77 THR THR THR ASP ASP ASP TRP	78 ALA VAL VAL SER THR SER 99 PHE PHE	YAL ALA 79 TYR TYR 88 ALA ALA ALA 100 ASP ASP	THR THR 80 MET MET MET VAL VAL 100A SER SER	81 GLU GLN 90 TYR TYR TYR TYR ASP	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GIY	82A SER SER SER CYS CYS CYS CYS 100D ARG
B C A B C A B C A	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C UEU LEU LEU 94 ARG ARG ARG	14 SER SER SER SER ARG ARG GIN GIN	PHE 75 SER*: THR THR 84 SER: SER SER 96 GLY GLY	76 FHR. SER SER SER GLU GLU 97 LEU 103	THR 77 THR THR THR ASP ASP ASP TRP 104	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE 105	VAL ALA 79 TYR TYR 88 ALA ALA ALA 100 ASP ASP 106	THR THR 80 MET MET MET VAL VAL 100A SER SER 107 THR	81 GLU GLN 90 TYR TYR TYR TYR ASP	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GIY	82A SER SER SER CYS CYS CYS CYS 100D ARG
B C A B C A B C A B	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C UEU LEU LEU 94 ARG ARG ARG 100F PEIE	14 SER SER SER SER ARG ARG ARG GLN GLN	PHE 75 SER*: THR THR 84 SER: SER SER 96 GLY GLY	76 FHR. SER SER SER GLU GLU 97 LEU 103-	THR 77 THR THR THR ASP ASP ASP TRP	78 ALA VAL VAL SER THR SER 99 PHE PHE	YAL ALA 79 TYR TYR 88 ALA ALA ALA 100 ASP ASP	THR THR 80 MET MET MET VAL VAL 100A SER SER	81 GLU GLN 90 TYR TYR TYR TYR 100B ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GIY 109	82A SER SER SER SER CYS CYS CYS LOOD ARG 110
B C A B C A B C A	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C UEU LEU LEU 94 ARG ARG ARG	14 SER SER SER SER ARG ARG GIN GIN 101 ASP	PHE 75 SER*: THR THR 84 SER: SER 96 GLY GLY 102 PHE	76 FHR. SER SER SER GLU GLU 97 LEU 103- TRP	THR 77 THR THR THR 86 ASP ASP ASP TRP 104 GLY	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE PHE 105 GLN GLN	VAL ALA 79 TYR TYR 88 ALA ALA ALA ALA 100 ASP ASP 106 GLY	THR THR 80 MET MET MET VAL VAL 100A SER 107 THR THR	81 GLU GLN 90 TYR TYR TYR 100B ASP ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GLY 109 VAL	82A SER SER SER CYS CYS CYS CYS 100D ARG 110 THR
B C A B C A B C A B	ASH 72 ASP ASP ASP SER SER SER SER THR ALA THR	GLU 73 GLU THR GLU 82C UEU LEU LEU 94 ARG ARG ARG 100F PEIE	14 SER SER SER SER ARG ARG GIN GIN 101 ASP	PHE 75 SER*: THR THR 84 SER: SER 96 GLY GLY 102 PHE	76 FHR. SER SER SER GLU GLU 97 LEU 103- TRP	THR 77 THR THR THR 86 ASP ASP ASP TRP 104 GLY	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE PHE 105 GLN GLN	VAL ALA 79 TYR TYR 88 ALA ALA ALA ALA 100 ASP ASP 106 GLY	THR THR 80 MET MET MET VAL VAL 100A SER 107 THR THR	81 GLU GLN 90 TYR TYR TYR 100B ASP ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GLY 109 VAL	82A SER SER SER CYS CYS CYS CYS 100D ARG 110 THR
B C A B C A B C	ASH 72 ASP ASP ASP SER SER SER SER 100E GLY	GLU 73 GLU THR GLU 82C LEU LEU 94 ARG ARG ARG PHE PHE	14 SER SER SER SER 83 TER ARG ARG CIN CLN 101 ASP ASP 113	PHE 75 SER*: THR THR 84 SER: SER 96 GLY GLY 102 PHE	76 FHR. SER SER SER GLU GLU 97 LEU 103- TRP	THR 77 THR THR THR 86 ASP ASP ASP TRP 104 GLY	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE PHE 105 GLN GLN	VAL ALA 79 TYR TYR 88 ALA ALA ALA ALA 100 ASP ASP 106 GLY	THR THR 80 MET MET MET VAL VAL 100A SER 107 THR THR	81 GLU GLN 90 TYR TYR TYR 100B ASP ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GLY 109 VAL	82A SER SER SER CYS CYS CYS CYS 100D ARG 110 THR
BC ABC ABC ABC A	ASH 72 ASP ASP ASP SER SER SER SER 100E GLY 111 VAL	GLU 73 GLU THR GLU 82C UEU LEU 94 ARG ARG ARG PHE 112 SER	14 SER SER SER SER 83 TER ARG ARG CIN CIN 101 ASP 113 SER	PHE 75 SER*: THR THR 84 SER: SER 96 GLY GLY 102 PHE	76 FHR. SER SER SER GLU GLU 97 LEU 103- TRP	THR 77 THR THR THR 86 ASP ASP ASP TRP 104 GLY	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE PHE 105 GLN GLN	VAL ALA 79 TYR TYR 88 ALA ALA ALA ALA 100 ASP ASP 106 GLY	THR THR 80 MET MET MET VAL VAL 100A SER 107 THR THR	81 GLU GLN 90 TYR TYR TYR 100B ASP ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GLY 109 VAL	82A SER SER SER CYS CYS CYS CYS 100D ARG 110 THR
B C A B C A B C	ASH 72 ASP ASP ASP SER SER SER SER 100E GLY	GLU 73 GLU THR GLU 82C LEU LEU 94 ARG ARG ARG PHE PHE	14 SER SER SER SER 83 TER ARG ARG CIN CLN 101 ASP ASP 113	PHE 75 SER*: THR THR 84 SER: SER 96 GLY GLY 102 PHE	76 FHR. SER SER SER GLU GLU 97 LEU 103- TRP	THR 77 THR THR THR 86 ASP ASP ASP TRP 104 GLY	ARG LYS 78 ALA VAL VAL 87 SER THR SER 99 PHE PHE 105 GLN GLN	VAL ALA 79 TYR TYR 88 ALA ALA ALA ALA 100 ASP ASP 106 GLY	THR THR 80 MET MET MET VAL VAL 100A SER 107 THR THR	81 GLU GLN 90 TYR TYR TYR 100B ASP ASP 108	THR THR 82 LEU LEU 91 TYR TYR TYR 100C GLY 109 VAL	82A SER SER SER CYS CYS CYS CYS 100D ARG 110 THR

	FIGU	RE 3· Y	VARIA	BLE RI	EGION	OF TH	E LIGH	IT CHA	AIN OF	IOR-R	3.	
	1	2	3 .	4	5 .	6	7	8	9	10	11	12
Α	ASP	VAL	LEU	MET	THR	GLN	ILE	PRO	LEU	SER SER	LEU	PRO
В	ASP	VAL	VAL	MET	THR	GLN GLN	SER ILE	PRO PRO	LEU LEU	SER	LEU LEU	PRO PRO
С	ASP	<u>VAL</u>	LEU	MET	THR	GLN	ш	rko	LLU	QL4C	220	110
	13	14	15	16	17	18	19	20	21	22 ·	23	24
A	VAL	SER	LEU	GLY	ASP	GLN	ALA	SER	ILE	SER	CYS	ARG
В	VAL	THR	LEU	GLY	GLN	PRO	ALA	SER	ILE	SER	CYS	400
С	VAL	SER	LEU	GLY	ASP	GLN	ALA	SER	ILE	SER	CYS	ARG
	25	26.	27	27A	27B	27C .	27D	27E	28	29	30	31
Α	SER	SEB	GLN	ASH	ILE	VAL .	His	SER	asn	GLY	ASH	THR
В									801		AON	. =0.5
С	SER	SER	GLM	ASH	ILE	VAL	HIS	SER	asn	GTA	ASN	THR
	32	33	34 .	35	36	37	38	39	40	41	42	43
À	TYR .	LEO	ASP	TRP	TYR	LEU	GLN	LYS	PRO	GLY	GLN	SER
В				TRP	PHE	GLN	GLN	ARG	PRO	GLY	GLN	SER
С	TYR	TEB	ASP	TRP	TYR .	LEU	GLN	LYS	PRO	GLY	GLN	SER
	44	45	46	47	48	49	50	51	52	53	54	55
A	PRO	ASN	LEU	LEU	ILE	TYR	LYS	VAL	SLA	ASH	ARC	PHE
В	PRO	ARG	ARG	LEU	ILE	TYR						130 120 110 110
С	PRO	ASN	LEU	LEU	ILE	TYR	LYS	VAL	SER	ASN	ARG	PHE
		710.1	220		. ===							
	56	57	58	59	60	61	62	63	64	65	66	67
A	56	57	58	59	60	61 ARG	62 PHE	63 ARG	64 GLY	65 SER	<u>GLY</u>	SER
A B	56	57 GLY GLY	58 VAL VAL	59 PRO PRO	60 ASP ASP	61 ARG ARG	62 PHE PHE	63 ARG SER	64 GLY GLY	65 SER SER	GLY GLY	SER SER
	56	57 GLY	58 VAL	59 PRO	60 ASP	61 ARG	62 PHE	63 ARG	64 GLY	65 SER	<u>GLY</u>	SER
В	56 SEB	57 GLY GLY GLY	58 VAL VAL VAL	59 PRO PRO PRO	60 ASP ASP	61 ARG ARG	62 PHE PHE	63 ARG SER	64 GLY GLY	65 SER SER	GLY GLY	SER SER
B C	56 SEB SEB 68	57 GLY GLY	58 VAL VAL	59 PRO PRO	60 ASP ASP ASP	61 ARG ARG ARG 73 LEU	62 PHE PHE PHE 74 LYS	63 ARG SER ARG 75 ILE	GLY GLY GLY SER	65 SER SER SER 77 ARG	GLY GLY GLY 78 VAL	SER SER SER 79 GLU
B C A B	56 SEB	57 GLY GLY GLY 69	58 VAL VAL VAL 70 ASP ASP	59 PRO PRO PRO 71 PHE PHE	60 ASP ASP ASP 72 THR THR	61 ARG ARG ARG 73 LEU LEU	62 PHE PHE PHE 74 LYS LYS	63 ARG SER ARG 75 ILE ILE	GLY GLY GLY GLY 76 SER SER	65 SER SER SER 77 ARG ARG	GLY GLY GLY 78 VAL VAL	SER SER SER 79 GLU GLU
B C	56 SEB SEB 68 GLY	57 GLY GLY GLY 69 THR	58 VAL VAL VAL 70 ASP	59 PRO PRO PRO 71 PHE	60 ASP ASP ASP 72 THR	61 ARG ARG ARG 73 LEU	62 PHE PHE PHE 74 LYS	63 ARG SER ARG 75 ILE	GLY GLY GLY SER	65 SER SER SER 77 ARG	GLY GLY GLY 78 VAL	SER SER SER 79 GLU
B C A B	56 SEB SEB 68 GLY GLY	57 GLY GLY GLY 69 THR THR	58 VAL VAL VAL 70 ASP ASP	59 PRO PRO PRO 71 PHE PHE	60 ASP ASP ASP 72 THR THR	61 ARG ARG ARG 73 LEU LEU	62 PHE PHE PHE 74 LYS LYS	63 ARG SER ARG 75 ILE ILE	GLY GLY GLY GLY 76 SER SER	65 SER SER SER 77 ARG ARG	GLY GLY GLY 78 VAL VAL	SER SER SER 79 GLU GLU
B C A B C	56 SEB 68 GLY GLY GLY	57 GLY GLY 69 THR THR THR	58 VAL VAL VAL 70 ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP 72 THR THR THR S4 GLY	ARG ARG ARG T3 LEU LEU LEU	PHE PHE PHE LYS LYS	63 APG SER ARG 75 ILE ILE ILE 87 TYR	GLY GLY GLY 76 SER SER SER SER CYS	65 SER SER SER 77 ARG ARG ARG	GLY GLY GLY 78 VAL VAL VAL	SER SER SER 79 GLU GLU GLU
B C A B	56 SEB 68 GLY GLY GLY 80	57 GLY GLY 69 THR THR THR	VAL VAL 70 ASP ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP 72 THR THR THR GLY GLY	ARG ARG ARG ARG - 73 LEU LEU LEU VAL VAL	62 PHE PHE PHE 74 LYS LYS LYS LYS TYR	63 SER ARG 75 ILE ILE ILE TYR	GLY GLY GLY SER SER SER CYS CYS	65 SER SER SER 77 ARG ARG ARG PHE	GLY GLY GLY 78 VAL VAL VAL 90 GL	SER SER SER 79 GLU GLU GLU 91 TVB
B C A B C	56 SEB 68 GLY GLY GLY ALA	57 GLY GLY 69 THR THR THR GLU	VAL VAL 70 ASP ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP 72 THR THR THR S4 GLY	ARG ARG ARG ARG - 73 LEU LEU LEU VAL	62 PHE PHE PHE 74 LYS LYS LYS LYS	63 APG SER ARG 75 ILE ILE ILE 87 TYR	GLY GLY GLY SER SER SER CYS CYS	65 SER SER SER 77 ARG ARG ARG	GLY GLY GLY 78 VAL VAL VAL 90	SER SER SER 79 GLU GLU GLU 91
B C A B C	SEB 68 GLY GLY GLY ALA ALA ALA	GLY GLY 69 THR THR THR GLU GLU	VAL VAL 70 ASP ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP 72 THR THR THR GLY GLY	ARG ARG ARG ARG - 73 LEU LEU LEU VAL VAL	62 PHE PHE PHE 74 LYS LYS LYS LYS TYR	63 SER ARG 75 ILE ILE ILE TYR	GLY GLY GLY SER SER SER CYS CYS	65 SER SER SER 77 ARG ARG ARG PHE	GLY GLY GLY 78 VAL VAL VAL 90 GL	SER SER SER 79 GLU GLU GLU 91 TVB
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B C A B C	56 SEB 68 GLY GLY GLY ALA ALA ALA 92	GLY GLY 69 THR THR THR GLU GLU GLU 93	VAL VAL 70 ASP ASP ASP ASP ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP 72 THR THR THR GLY GLY GLY GLY TRP	ARG ARG ARG ARG 73 LEU LEU LEU VAL VAL 97 THB	62 PHE PHE PHE T4 LYS LYS LYS LYS TYR TYR TYR PHE PHE	63 AFG SER ARG 75 ILE ILE ILE 87 TYR TYR TYR TYR GLY GLY	GLY GLY GLY 76 SER SER SER CYS CYS CYS CYS GLY GLY	65 SER SER SER 77 ARG ARG ARG PHE 101 GLY GLY	GLY GLY GLY 78 VAL VAL VAL 90 GLII 102 THR THR	SER SER SER 79 GLU GLU 91 TVB 103 LYS LYS
B C A B C	56 SEB 68 GLY GLY GLY ALA ALA ALA 92	GLY GLY 69 THR THR THR GLU GLU GLU 93	VAL VAL 70 ASP ASP ASP ASP ASP ASP	PROPROPROPROPROPROPROPROPROPROPROPROPROP	ASP ASP ASP THR THR THR GLY GLY GLY GLY	ARG ARG ARG ARG 1EU LEU LEU LEU VAL VAL 97	62 PHE PHE PHE T4 LYS LYS LYS LYS TYR TYR TYR PHE	63 AFG SER ARG 75 ILE ILE ILE 87 TYR TYR TYR TYR 99 GLY	GLY GLY GLY 76 SER SER SER CYS CYS CYS CYS GLY	65 SER SER SER 77 ARG ARG ARG PHE 101 GLY	GLY GLY GLY 78 VAL VAL VAL 90 GLII 102 THR	SER SER SER 79 GLU GLU GLU 91 TVB
B C A B C	SEB 68 GLY GLY GLY SLA ALA ALA 92 SEB	GLY GLY 69 THR THR THR GLU GLU GLU 93 HIS	VAL VAL 70 ASP	PROPROPRO 71 PHE PHE PHE VAL LEU 95 PRO PRO	ASP ASP ASP 72 THR THR THR GLY GLY GLY GLY TRP	ARG ARG ARG ARG 73 LEU LEU LEU VAL VAL 97 THB	62 PHE PHE PHE T4 LYS LYS LYS LYS TYR TYR TYR PHE PHE	63 AFG SER ARG 75 ILE ILE ILE 87 TYR TYR TYR TYR GLY GLY	GLY GLY GLY 76 SER SER SER CYS CYS CYS CYS GLY GLY	65 SER SER SER 77 ARG ARG ARG PHE 101 GLY GLY	GLY GLY GLY 78 VAL VAL VAL 90 GLII 102 THR THR	SER SER SER 79 GLU GLU 91 TVB 103 LYS LYS
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B C A B C	56 SEB 68 GLY GLY GLY SLA ALA ALA ALA 92 SEB 5EB	GLY GLY 69 THR THR THR GLU GLU GLU 93 HIS HIS	VAL VAL 70 ASP	59 PRO PRO PRO 71 PHE PHE PHE VAL LEU 95 PRO PRO PRO	ASP ASP ASP 72 THR THR THR GLY GLY GLY TRP 108	ARG ARG ARG ARG 73 LEU LEU LEU VAL VAL 97 THB	62 PHE PHE PHE T4 LYS LYS LYS LYS TYR TYR TYR PHE PHE	63 AFG SER ARG 75 ILE ILE ILE 87 TYR TYR TYR TYR GLY GLY	GLY GLY GLY 76 SER SER SER CYS CYS CYS CYS GLY GLY	65 SER SER SER 77 ARG ARG ARG PHE 101 GLY GLY	GLY GLY GLY 78 VAL VAL VAL 90 GLII 102 THR THR	SER SER SER 79 GLU GLU 91 TVB 103 LYS LYS







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C A B	SER 60 SER	61	62	63	64	65	66 ARG	67 PHE	68 THR	69 ILE	70 SER	71 <u>ARG</u>
C A	SER 60	61 ARG	62 GLN	63 CYS CYS	64 GLU	GLY GLY	66 ARG ARG ARG	67 PHE PHE PHE	68 THR THR THR	69 ILE ILE ILE	70 SER SER SER	71 ARG ARG ARG
C A B	SER 60 SER	61 ARG	62 GLN GLN	63 CYS CYS	64 GLU GLU 76	65 GLY GLY 77	66 ARG ARG ARG	67 PHE PHE PHE	68 THR THR THR 80	69 ILE ILE ILE	70 SER SER SER	71 ARG ARG ARG 82A
C A B	SER 60 SIB SIB 72 ASP	61 ARG ARG 73 ASN	62 GIN GIN 74 VAL	63 CYS CYS 75 LYS	64 GLU GLU 76 ASN	65 GLY CLY 77 THIR	66 ARG ARG ARG 78 LEU	67 PHE PHE PHE 79 TYR	68 THR THR THR THR 80 LEU	69 ILE ILE ILE GLN	70 SER SER SER SER MET	71 ARG ARG ARG 82A SER
C A B C A B	SER 60 SIB SEB 72 ASP ASP	61 ARG ARG 73 ASN ASN	62 GIN GIN 74 VAL SER	63 CYS CYS LYS LYS	64 GLU GLU 76 ASN ASN	65 GLY GLY 77 THIR THIR	66 ARG ARG ARG 78 LEU LEU	67 PHE PHE PHE 79 TYR TYR	68 THR THR THR LEU LEU	69 ILE ILE ILE GLN GLN	70 SER SER SER SER MET MET	71 ARG ARG ARG 82A SER ASN
C A B C	SER 60 SIB SIB 72 ASP	61 ARG ARG 73 ASN	62 GIN GIN 74 VAL	63 CYS CYS 75 LYS	64 GLU GLU 76 ASN	65 GLY CLY 77 THIR	66 ARG ARG ARG 78 LEU	67 PHE PHE PHE 79 TYR	68 THR THR THR THR 80 LEU	69 ILE ILE ILE GLN	70 SER SER SER SER MET	71 ARG ARG ARG 82A SER
C A B C A B	SER 60 SIB 5EB 72 ASP ASP ASP	ARG ARG 73 ASN ASN	62 GIN GIN 74 VAL SER VAL	63 CYS CYS 175 LYS LYS LYS	64 GLU GLU 76 ASN ASN	65 GLY GLY 77 THIR THIR THIR	66 ARG ARG ARG 78 LEU LEU LEU	67 PHE PHE PHE 79 TYR TYR TYR	68 THR THR THR 80 LEU LEU LEU	69 ILE ILE ILE 81 GLN GLN GLN	70 SER SER SER 82 MET MET MET	71 ARG ARG ARG SER ASN SER
C A B C	SER 60 SIB 72 ASP ASP ASP ASP	61 ARG ARG 73 ASN ASN ASN 82C	62 GIN GIN 74 VAL SER VAL 83	63 CYS CYS 75 LYS LYS LYS	64 GLU GLU 76 ASN ASN ASN	65 GLY GLY 77 THR THR THR THR	66 ARG ARG ARG 78 LEU LEU LEU	67 PHE PHE PHE 79 TYR TYR TYR	68 THR THR THR 80 LEU LEU LEU	69 ILE ILE ILE GLN GLN GLN 90	70 SER SER SER SER MET MET MET	71 ARG ARG ARG SER ASN SER
C A B C A B C	SER 60 SIB 72 ASP ASP ASP ASP SER	61 ARG ABG 73 ASN ASN ASN ASN	62 GLN 74 VAL SER VAL 83 ARG	63 CYS CYS 75 LYS LYS LYS LYS	64 GLU 76 ASN ASN ASN SS GLU	65 GLY CLY 77 THR THR THR THR ASP	66 ARG ARG ARG 78 LEU LEU LEU LEU	67 PHE PHE PHE 79 TYR TYR TYR TYR	68 THR THR THR 80 LEU LEU LEU MET	69 ILE ILE ILE ILE 81 GLN GLN GLN TYR	70 SER SER SER 82 MET MET MET 91 TYR	71 ARG ARG ARG SER ASN SER
C A B C A B	SER 60 SIB 72 ASP ASP ASP SER SER SER	61 ARG ABG 73 ASN ASN ASN E2C LEU LEU	62 GLN 74 VAL SER VAL 83 ARG ARG	63 CYS CYS 75 LYS LYS LYS LYS SER ALA	64 GLU 76 ASN ASN ASN GLU GLU	65 GLY 77 THR THR THR THR ASP ASP	66 ARG ARG ARG 78 LEU LEU LEU THR THR	67 PHE PHE PHE 79 TYR TYR TYR TYR	68 THR THR THR 80 LEU LEU LEU LEU MET VAL	69 ILE ILE ILE 81 GLN GLN GLN TYR TYR	70 SER SER SER SER MET MET MET	71 ARG ARG ARG SER ASN SER 92 CYS
C A B C A B C	SER 60 SIB 72 ASP ASP ASP ASP SER	61 ARG ABG 73 ASN ASN ASN ASN	62 GLN 74 VAL SER VAL 83 ARG	63 CYS CYS 75 LYS LYS LYS LYS	64 GLU 76 ASN ASN ASN SS GLU	65 GLY CLY 77 THR THR THR THR ASP	66 ARG ARG ARG 78 LEU LEU LEU LEU	67 PHE PHE PHE 79 TYR TYR TYR TYR	68 THR THR THR 80 LEU LEU LEU MET	69 ILE ILE ILE ILE 81 GLN GLN GLN TYR	70 SER SER SER 82 MET MET MET TYR TYR	71 ARG ARG ARG SER ASN SER 92 CYS CYS
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C A B C A B C	SER 60 SIB 72 ASP ASP ASP ASP SER SER SER SER SER 93	61 ARG ARG 73 ASN ASN ASN E2C LEU LEU LEU	62 GLN 74 VAL SER VAL 83 ARG ARG ARG	63 CYS CYS LYS LYS LYS LYS SER ALA SER	64 GLU 76 ASN ASN ASN GLU GLU GLU	65 GLY 77 THR THR THR ASP ASP 98	66 ARG ARG ARG 78 LEU LEU LEU THR THR THR	67 PHE PHE PHE 79 TYR TYR TYR TYR ALA ALA ALA	68 THR THR THR 80 LEU LEU LEU LEU MET VAL MET	69 ILE ILE ILE SI GLN GLN GLN TYR TYR TYR	70 SER SER SER 82 MET MET MET TYR TYR TYR TYR	71 ARG ARG ARG SER ASN SER 92 CYS CYS CYS
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В	TYR								GLY	ILE	PRO	ALA
С	<u>TYR</u>	SEB	ALA	SER	SEA	ARB	ASH	SER	GLY	VAL	PRO	ALA
	61	62	63	64.	65	66	67	68	69	70	71	72
Α	ARG		THR	GLY:			SER	GLY	THR	ASP	PHE	THR
В	ARG	PHE	SER	GLY	SER	<u>GLY</u>	SER	GLY	THR	GLU	PHE	THR
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			75	76	77	78	79	80	81	82	83	84
A	73 LEU	74 THR	ILE	SER	ASN	VAL	GIN	SER	GU	ASP	SLEU!	ALA
В	LEU	THR	ILE	SER	ARG	LEU	GLN	SER	GLU	ASP	PHE	ALA
č	LEU	THR	ILE	SER	ASN	VAL	GLN	SER	GLU	ASP	PHE	ALA
		0.6	07		89	90	91	92	93	94	95	95A
A	85 GLU	86 TYR	87 PHE	. 88 CYS	GLH	ELI		ASN	SIR	178		TO T
A B	VAL	TYR	TYR	CYS			3		SOCIETATION.	OO BUT ALL DE WOOD	062.10220000	
c	VAL	TYR	TYR	CYS	CTM	CLN	TYB	ash	SEB	TYB	PRO	LEO
			00		100	101	102	103	104	105	106	107
	96	97 Tub	98 <u>PHE</u>	99 GLY	100 ALA	101 . GLY		LYS	LEU	GLU		LYS
А В ··	YAL	IMM	PHE	GLY	GLN	GLY	THR	ARG	VAL	GLU	ILE	LYS
C	VAL	THA	PHE	GLY	ALA	GLY	THR	LYS	LEU	GLU	LEU	LYS
		100										
	108 ARG	109 ALA										
A B	ARG	GLU										
Č	ARG	ALA						•				

	FIGUR	E 9: VA	ARIABI	E REC	O MOI	F THE	HEAV	Y CHA	IN OF I	OR-CE	A-1.	
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Α	POEN.	PRO	LYS									VAL
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			.:				10	20		22	• •	
	13	14	15 GLU	16 ALA	. 17	18 LEU	19 ASN	20 CYS	21 SER	22 CYS	23 ALA	24 VAL
A	LYS	PRO	GLY	ALA	SER SER	LEU	LYS	VAL	SER	CYS	LYS	ALA
В	LYS			ALA		LEU	ASN	CYS	SER	CYS	ALA	VAL .
С	LYS	PRO	GLY	ALA	SER	LEU	W2IA	CIS	SER	CIS	ALA	VAL
	25	26	27	28	29	30	31	32	33	34 -	35	36
Α	SER	GLY	PHE	PRO	PHE		ARB	TYB	ALA	MET	SER	TRP
В	SER	GLY	TYR	THR	PHE	THR						TRP
Ċ.	SER	GLY	PHE	PRO	PHE	ASN	ARG	TYB	ALA	MET	SEB	TRP
		. —										
	37	· 38	39	40	41	42	43	44	45	46	47	48 -
A.	VAL	LEU	GLN	THR	PRO	GLU	LYS		LEU	GLU	TRP	VAL
В	VAL	ARG	GLN	ALA	PRO	GLY	GLN	ARG	LEU	GLU	TRP	MET
C	VAL	LEU	GĽŅ	THR	PRO	GLU	LYS	ARG	LEU	GLU	TRP	VAL
	49	50	51	52	52A	53	54	55	56	57	58	59
A	ALA	PHE	ILE	SEB	SER	ASP	ASP	GLY	ILE	ALA	TYR	TYB
В	GLY				,		-					
C	ALA ·	PHE	ILE	SEA	SER	ASP	ASP	ELY	ILE	ALA	TYB	TYB
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	60	61	62	63	64	65	66	67	68	69	70	71
A	ALA	810	SEA	LYS	GIA	TYB	ARG	PHE	THR	ILE	SER	ARG
_B							ARG	VAL	THR	ILE	THR	ARG
С	AIA	elv	SER	LYS	GLY	TYB	ARG	PHE	THR	ILE	SER	ARG
	72	73	74	75	76	77	78	79	80	81	82	82A
Α	ASP	ASN	ALA	LYS	ASN	ILE	LEU	TYR	LEU	GLN	MET	SER
В	ASP	THR	SER	ALA	SER	THR	ALA	TYR	MET	GLU	· LEU	SER
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	82B	82C	83	84	85	86	87	88	89	90	91	92 .
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B C	SER	LEU	ARG	SER	GLN	ASP ASP	THR THR	ALA ALA	VAL	TYR	TYR	CYS
	JLK	LLC	ANO	JLK	OLIV	AUI	ш	, ALA	VAL.	111	· IK	
	93	94	95	96	97	98	99	100	100A	100B	100C	101
A	ALA	ARG	VAL	JyB	T/B	TYB	GLY	SIR	31	TIE		169
В	ALA	ARG										
C ·	ALA	ARG	AVT	TYR	TYB	TYR	GLY	SEA	SER	777	Phe	ASP
Α	102	103	104	105	106	107	108	109	110	111	112	113
В	TYR	inp!		GIN			TION				SER	SER
Č	Secretary of the Secretary of	TRP	GLY	ĞLÜ	GLY	THR	LEU	VAL	THR	VAL	SER	SER
-	TYR	TRP	GLY	GLN	GLY	THR	LEU	VAL	THR	VAL	SER	SER
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	FIG	IRE 10)·VARI	ABLE	REGIO	N OF I	HE LI	GHT O	FIOR-C	CEA 1.		
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Α	ASP	TE	GEN			OEN	SPR	PRO	CER	THR	LEU	SER.
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С	ASP	ILE .	<u>GLN</u>	MET	THR	GLN	SER	PRO	SER	IAK .	LEU	SER .
					17	18	19	20	21	22	23	24
	13	14 SER	15 VAL	16 GLY	ASP	ARG	VAL	SER	VAL	THR	CYS	LYS
A	Process of the second	SER	VAL	GLY	ASP	SER	ILE	THE	ILE	THR	CYS	•
В	ALA	SER	VAL	GLY	ASP	ARG	VAL	SER	VAL	THR	CYS	LYS
С	ALA	SER	VAL	02.	,							•
	25	26	27	28	29	30	31	32	33	34	35 .	36
A	ALA	SIB	GLH	ASH	ALA	GLY	ILE	ASH	VAL	ALA	TRP	TYR
В					•						TRP	PHE
č	ALA	SER	GLN	ASH	ALA	CLY	ILE	ash	VAL	ALA	TRP ·	TYR
_									45	46 .	47	48
	37	38	39	40	41	42	43 CED	44 PRO	LYS	ALA	LEU	ILE
A	GLN	GLN	LYS	PRO	GLY GLY	GLN LYS	SER ALA	PRO	ASN	VAL	LEU	ILE
В	GLN	GLN	LYS	PRO PRO	GLY	GLN	SER	PRO	LYS	ALA	LEU	ILE
С	GLN	GLN	LYS	PRO	OD I	OLA V	000					
	49	50	51	52	53	54	55	56	57	58	59	60
Α	TYR	SER	ALA	SEB	SEB	ARG	ASH	लंडा)				ASP
В	<u>TYR</u>		•						GLY	ILE	PRO	SER
С	TYR	SER	ALA	SER	SEB	ARG	ash	SER	GLY	ILE	PRO	SER
	4		1				49	40	60	70	71	72
	61	62	63	64	65 65	66	67 	68	69	70 .	71 PHE	72 THR
A	ARO	្សារូវភ	on in	(GIBY	SER	(RE)	Mar	OLY!	OHR	70 -ASP GLU		72 THR THR
В	ARG	PHE	THR. SER	GLY GLY	SER SER	GLY GLY	67 SER SER SER	OLY GLY		4/4SPL/	PHE	THR
	ARO	្សារូវភ	on in	(GIBY	SER	(RE)	SER SER	OLY!	THR	GLU	PHE	THR THR THR
В	ARG	PHE PHE	SER SER	GLY GLY	SER SER SER	GLY GLY GLY	SER SER SER	GLY GLY	THR THR THR	GLU GLU	PHE PHE PHE	THR THR THR
В	ARG ARG ARG 73 LEU	PHE PHE	SER SER 75	GLY GLY GLY 76 SER	SER SER SER 71	GLY GLY GLY 78	SER SER SER 79	GLY GLY GLY 80 SERV	THR THR THR 81	GLU GLU 82 ASI	PHE PHE PHE 83	THR THR THR 84
B C A B	ARG ARG ARG 73 LEU LEU	PHE PHE PHE 74 THR	SER SER 75 USA	GLY GLY GLY 76 SER ILE	SER SER SER 77 77 ASN	GLY GLY GLY 78 VAL LEU	SER SER SER 79 GLN	GLY GLY GLY 80 SER SER	THR THR THR 81 GDQ ASP	GLU GLU 82 ASP	PHE PHE PHE 83 PHE	THR THR THR 84 ALA
B C	ARG ARG ARG 73 LEU	PHE PHE PHE 74	SER SER 75	GLY GLY GLY 76 SER	SER SER SER 71	GLY GLY GLY 78	SER SER SER 79	GLY GLY GLY 80 SERV	THR THR THR 81	GLU GLU 82 ASI	PHE PHE PHE 83	THR THR THR 84
B C A B	ARG ARG ARG 73 LEU LEU	PHE PHE PHE 74 THR	SER SER 75 USA	GLY GLY 76 SER ILE	SER SER SER 77 ASN ASN	GLY GLY 78 LEU LEU	SER SER SER 79 GLN GLN GLN	GLY GLY BO SER SER SER	THR THR 81 GIN ASP ASP	GLU GLU 82 ASP ASP	PHE PHE 83 PHE PHE	THR THR THR ALA ALA
B C A B C	ARG ARG ARG 73 LEU LEU LEU	PHE PHE PHE THR THR	SER SER 75 VAL VAL 87	GLY GLY 76 SER ILE ILE	SEB SER SER 77 3N ASN ASN B9	GLY GLY 78 LEU LEU	SER SER SER 79 GLN GLN 91	GLY GLY 80 SFR SER SER 92	THR THR 81 GING ASP ASP	GLU GLU 82 ASP ASP ASP 94	PHE PHE 83 PHE PHE PHE	THR THR THR ALA ALA 95A
B C A B C	ARG ARG ARG 73 LEU LEU LEU	PHE PHE 74 PHR THR THR	SER SER 75 TUE VAL VAL 87	GLY GLY 76 SER ILE ILE 88	SEB SER SER 77 3N ASN ASN B9	GLY GLY 78 LEU LEU	SER SER SER 79 GLN GLN 91	GLY GLY 80 SFR SER SER 92	THR THR 81 GING ASP ASP	GLU GLU 82 ASP ASP ASP 94	PHE PHE 83 PHE PHE PHE	THR THR THR ALA ALA
B C A B C	ARG ARG ARG 73 LEU LEU LEU LEU THR	PHE PHE 74 THR THR TYR	THE SER SER 75 THE VAL VAL VAL 87 PHE TYR	GLY GLY 76 SER ILE ILE 88 CYS	SER SER SER 77 ASN ASN ASN 89	GLY GLY 78 VAL LEU LEU 90	SER SER SER 79 GLN GLN GLN 91	GLY GLY GLY 80 SER SER SER 92	THR THR 81 GUN ASP ASP 93	ASP GLU GLU 82 ASP ASP ASP 94	PHE PHE 83 24 W PHE PHE 95 1 A D S	THR THR THR 84 ALA ALA 95A
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B C A B C	ARG ARG ARG 73 LEU LEU LEU THR	PHE PHE 74 THR THR TYR TYR	SER SER 75 VAL VAL 87 PEIS TYR TYR	GLY GLY GLY 76 SER ILE ILE CYS CYS	SER SER SER 77 ASN ASN ASN B9 GUN	GLY GLY 78 AL LEU LEU 90 GLY	SER SER SER 79 GLN GLN 91 TVB	GLY GLY 80 SER SER 92 ASN	THR THR 81 GUN ASP ASP 93 SUD	ASP GLU GLU 82 ASP ASP ASP 94 TYB	PHE PHE 83 PHE PHE PHE PHE PHE PHE PHE PHE	THR THR THR 84 ALA ALA 95A
B C A B C	ARG ARG ARG 73 LEU LEU LEU THR THR	PHE PHE 74 THR THR TYR TYR TYR	SER SER 75 UE VAL VAL 87 PPIE TYR TYR 98	GLY GLY 76 SER ILE ILE 88 CYS CYS 99	SER SER SER 77 SM ASN ASN B9 GH	GLY GLY 78 VAL LEU LEU 90 GUY	SER SER SER 79 GLN GLN 91 TVB	GLY GLY GLY 80 SER SER SER 92	THR THR 81 GUN ASP ASP 93	ASP GLU GLU 82 ASP ASP ASP 94	PHE PHE 83 24 W PHE PHE 95 1 A D S	THR THR THR 84 ALA ALA 95A
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B C A B C A B	ARG ARG ARG 73 LEU LEU LEU THR THR YAL	PHE PHE 74 THR THR TYR TYR 97	THE SER SER 75 HE VAL VAL 87 PHE PHE	GLY GLY GLY 76 SER ILE ILE CYS CYS CYS GLY	SER SER SER 77 ASN ASN ASN B9 GUN 100 AIA	GLY GLY 78 LEU LEU 90 GLY GLY GLY GLY GLY	SER SER SER GLN GLN WB 102 THR THR	GLY GLY 80 SER SER 92 ASN 103	THR THR 81 GUN ASP ASP 93 SUD SUB	ASP GLU GLU 82 ASP ASP ASP 94 TYB 105 GLN	PHE	THR THR THR 84 ALA ALA 95A LEU 107 LYS
B C A B C A B C	ARG ARG ARG 73 LEU LEU LEU THR THR	PHE PHE 74 THR THR TYR TYR TYR	THR SER SER 75 HE VAL VAL 87 PHE TYR TYR 98	GLY GLY GLY 76 SER ILE ILE CYS CYS CYS GLY	SER SER SER 77 ASN ASN ASN 6HH 100	GLY GLY 78 LEU LEU 90 GLY GLY GLY GLY GLY	SER SER SER GLN GLN WAS IO2 THR THR THR	GLY GLY 80 SER SER 92 ASN 103 LYS LYS	THR THR 81 GUN ASP ASP 93 SUD SUB UVAL	GLU GLU 82 ASP ASP ASP 105 GLN LEU	PHE	THR THR THR 84 ALA ALA 95A LEU 107 LYS LYS
B C A B C A B	ARG ARG ARG 73 LEU LEU LEU THR THR YAL	PHE PHE 74 THR THR TYR TYR 97	THE SER SER 75 HE VAL VAL 87 PHE PHE	GLY GLY GLY 76 SER ILE ILE CYS CYS CYS GLY	SER SER SER 77 ASN ASN ASN B9 GUN 100 AIA	GLY GLY 78 LEU LEU 90 GLY GLY GLY GLY GLY	SER SER SER GLN GLN WB 102 THR THR	GLY GLY 80 SER SER 92 ASN 103 LYS LYS	THR THR 81 GUN ASP ASP 93 SUD SUB UVAL	GLU GLU 82 ASP ASP ASP 105 GLN LEU	PHE	THR THR THR 84 ALA ALA 95A LEU 107 LYS LYS
B C A B C A B	ARG ARG ARG 73 LEU LEU LEU 85 OLU THR THR THR VAL	PHE PHE 74 THR THR TYR TYR TYR THB	THE SER SER 75 HE VAL VAL 87 PHE PHE	GLY GLY GLY 76 SER ILE ILE CYS CYS CYS GLY	SER SER SER 77 ASN ASN ASN B9 GUN 100 AIA	GLY GLY 78 LEU LEU 90 GLY GLY GLY GLY GLY	SER SER SER GLN GLN WAS IO2 THR THR THR	GLY GLY 80 SER SER 92 ASN 103 LYS LYS	THR THR 81 GUN ASP ASP 93 SUD SUB UVAL	GLU GLU 82 ASP ASP ASP 105 GLN LEU	PHE	THR THR THR 84 ALA ALA 95A LEU 107 LYS LYS
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MOUSE HEAVY CHAIRS FAMILY	1-TFS-139'CL	2-E7'CL	3-DFR-611.1'	171 74 145	2-MDC 450'F	6-TE2-36'PI	37 00:331-0	0-1/1 OFFW 1	2-12-0-12-0	10-H17-B5'6	13-36-11	12-T1-C	12-H17-78-C	14-14-14-14	10-11-11 10-11-11	13-0/ 55	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	7.6.00.71-11	10.96-75H-91	19-71 0 .45 CL	20-H37-24 CL	21-H37-42,CL	22-42.783.28	. 23-A10'CL	24-H37-88'CL	25-JD211CL	26-1 64-0	27-19/20-01	10 - 17 C - 0C	107-07 107-07	D TIMA-67	30-MS2.CL	31-VEMI'CL	32-WG1,CL	33-X-T151'CL

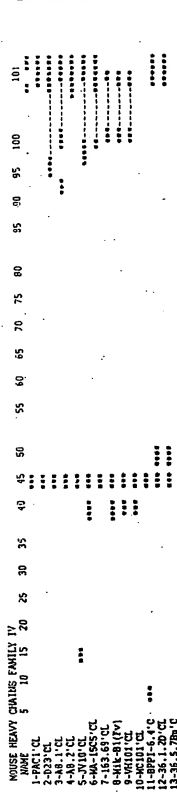
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	MOUSE HEAVY CHAINS FAMILY 11 NAME S 10 15 2	10.887-1	2-DF4-29.4'C	3-rab D'CL	4-BAT123'CL	5-35-20	6-Min2	7-ANG2'CL	R-40-120	9-40-40	10-H146-24F91	11-Mn7'CL	12-40-160	13-51 2-01	17-31-6	007-01-17	13-3/.1.1.CL	10-7-Finan-91	17-8-1-12-58	18-S27 CL	19-MM3-CL	20-111-141-01	21.MM1.01	77.1007-17	22-VHTF'CL	23-MOPC 315'C	24-MOPC315

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NOUSE HEAVY CHAIRS FAMILY V	2	145031	2-HOPC 104Z	3-MEX.	4-MPX25	ב השפתים	25.5	6-10CK	7-10EXS	8-10EX6	1220年		1-5004	12-3558	13- 七 0237	2 + 10 CC	12-FBEX12	6-414.2°C	7-262.9'6	8-126.33°CL	19-9.14.7°CL	0-Y0E(10	1-70EX14	22-16.3'CL	23-FDEX9	4-AC38 205.1	25-9.4.5'CL	26-4.1.3C4'CL	27-3.2.3ES'CL	28-45.21.1°CL	29-4.9.1'0	30-3.4.106'CL	31-42.504.28	32-4.311'CL	33-37.1E5.2a'	34-59.102.2'C	35-2,31,1'CL	36-MRAICHICL	17-165, 60'C	0.1 CH 31-0C	30-V2-016-1	27-80/ C2 C2: C1	40-163,72°CL	41-178.459.05	42-A20/44'CL	43-109-105.55	44-1.29°CL	45-L1.15'CL

Table 6

NOUSE HEAVY CHAINS FAMILY VI 1202.135°CL 2-202.61°CL 3-202.80°CL 4-202s.30°CL 6-111.109°CL 6-111.109°CL 9-AN11°CL 9-AN11°CL 9-AN11°CL 11-BAA 031 VH 11-BAA 031 VH																				***************************************	
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